

JAP20 Rec'd PCT/PTO 20 JUN 2006

2 - (HETERO) ARYL-SUBSTITUTED TETRAHYDROQUINOLINE DERIVATIVES

BACKGROUND OF THE INVENTION

The invention had the object of finding novel compounds having valuable properties, in particular those which can be used for the preparation of medicaments.

The present invention relates to compounds of the formula I and to the use thereof for the treatment and prophylaxis of diseases in which the inhibition, regulation and/or modulation of mitotic motor proteins, in particular the mitotic motor protein Eg5, plays a role, furthermore to pharmaceutical compositions which comprise these compounds.

In detail, the present invention relates to compounds of the formula I which preferably inhibit, regulate and/or modulate one or more mitotic motor proteins, to compositions which comprise these compounds, and to methods for the use thereof for the treatment of diseases and complaints such as angiogenesis, cancer, tumour formation, growth and propagation, arteriosclerosis, ocular diseases, choroidal neovascularisation and diabetic retinopathy, inflammatory diseases, arthritis, neurodegeneration, restenosis, wound healing or transplant rejection. In particular, the compounds according to the invention are suitable for the therapy or prophylaxis of cancer diseases.

During mitosis, various kinesins regulate the formation and dynamics of the spindle apparatus, which is responsible for correct and coordinated alignment and separation of the chromosomes. It has been observed that specific inhibition of a mitotic motor protein – Eg5 – results in collapse of the spindle fibres. The result of this is that the chromosomes can no longer be distributed correctly over the daughter cells. This results in mitotic arrest

and can thus cause cell death. Upregulation of the motor protein Eg5 has been described, for example, in tissue from breast lung and colon tumours. Since Eg5 takes on a mitosis-specific function, it is principally rapidly dividing cells and not fully differentiated cells that are affected by Eg5 inhibition. In addition, Eg5 regulates exclusively the movement of mitotic microtubuli (spindle apparatus) and not that of the cytoskeleton. This is crucial for the side-effect profile of the compounds according to the invention, since, for example, neuropathies, as observed in the case of Taxol, do not occur or only do so to a weakened extent. The inhibition of Eg5 by the compounds according to the invention is therefore a relevant therapy concept for the treatment of malignant tumours.

In general, all solid and non-solid tumours can be treated with the compounds of the formula I, such as, for example, monocytic leukaemia, brain, urogenital, lymphatic system, stomach, laryngeal and lung carcinoma, including lung adenocarcinoma and small-cell lung carcinoma. Further examples include prostate, pancreatic and breast carcinoma.

Surprisingly, it has been found that the compounds according to the invention effect specific inhibition of mitotic motor proteins, in particular Eg5. The compounds according to the invention preferably exhibit an advantageous biological activity which can easily be detected in the assays described herein, for example. In such assays, the compounds according to the invention preferably exhibit and cause an inhibiting effect, which is usually documented by IC₅₀ values in a suitable range, preferably in the micromolar range and more preferably in the nanomolar range.

As discussed herein, effects of the compound according to the invention are relevant to various diseases. Accordingly, the compounds according to the invention are useful in the prophylaxis and/or treatment of diseases which are influenced by inhibition of one or more mitotic motor proteins, in particular Eg5.

The present invention therefore relates to compounds according to the invention as medicaments and/or medicament active ingredients in the treatment and/or prophylaxis of the said diseases and to the use of compounds according to the invention for the preparation of a pharmaceutical for the treatment and/or prophylaxis of the said diseases, and also to a method for the treatment of the said diseases comprising the administration of one or more compounds according to the invention to a patient in need of such an administration.

It can be shown that the compounds according to the invention have an advantageous effect in a xenotransplant tumour model.

The host or patient can belong to any mammal species, for example a primate species, particularly humans; rodents, including mice, rats and hamsters; rabbits; horses, cattle, dogs, cats, etc. Animal models are of interest for experimental investigations, providing a model for the treatment of a human disease.

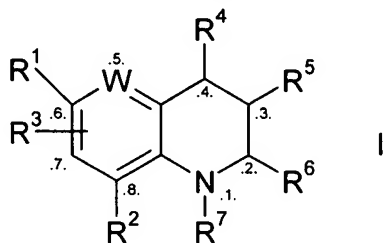
The susceptibility of a certain cell to treatment with the compounds according to the invention can be determined by testing in vitro. Typically, a culture of the cell is combined with a compound according to the invention at various concentrations for a period which is sufficient to enable the active ingredients to induce cell death or inhibit migration, usually between approximately one hour and one week. For testing in vitro, cultivated cells from a biopsy sample can be used. The viable cells remaining after the treatment are then counted.

The dose varies depending on the specific compound used, the specific disease, the patient status, etc. Typically, a therapeutic dose is sufficient considerably to reduce the undesired cell population in the target tissue, while the viability of the patient is maintained. The treatment is generally continued until a considerable reduction has occurred, for example at least

about a 50% reduction in the cell burden, and can be continued until essentially no undesired cells are detected in the body.

SUMMARY OF THE INVENTION

The invention relates to compounds of the formula I



in which

W denotes CH or N,

R¹, R², R³, independently of one another, denote H, A, aryl, heteroaryl, Hal, -(CY₂)_n-SA, -(CY₂)_n-SCF₃, -(CY₂)_n-SCN, -(CY₂)_n-CF₃, -(CY₂)_n-OCF₃, cycloalkyl, -SCH₃, -SCN, -CF₃, -OCF₃, -OA, -(CY₂)_n-OH, -(CY₂)_n-CO₂R, -(CY₂)_n-CN, -(CY₂)_n-Hal, -(CY₂)_n-NR₂, (CY₂)_n-OA, (CY₂)_n-OCOA, -SCF₃, (CY₂)_n-CONR₂, -(CY₂)_n-NHCOA, -(CY₂)_n-NHSO₂A, SF₅, Si(CH₃)₃, CO-(CY₂)_n-CH₃, -(CY₂)_n-N-pyrrolidone, CH(CH₂)_nNRCOOR, CHNRCOOR, NCO, CH(CH₂)_nCOOR, NCOOR, CH(CH₂)_nOH, N(CH₂)_nOH, CHNH₂, CH(CH₂)_nNR₂, CH(CH₂)_nNR₂, C(OH)R, CHNCOR, CH(CH₂)_n-aryl, CH(CH₂)_n-heteroaryl, CH(CH₂)_nR¹, N(CH₂)_nCOOR, CH(CH₂)_nX(CH₂)_n-aryl, CH(CH₂)_nX(CH₂)_n-heteroaryl, N(CH₂)_nCONR₂, XCONR(CH₂)_nNR₂, N[(CH₂)_nXCOOR]CO(CH₂)_n-aryl, N[(CH₂)_nXR]CO(CH₂)_n-aryl, N[(CH₂)_nXR]CO(CH₂)_nX-aryl, N[(CH₂)_nXR]SO₂(CH₂)_n-aryl, N[(CH₂)_nNRCOOR]CO(CH₂)_n-aryl, N[(CH₂)_nNR₂]CO(CH₂)_n-

aryl, N[(CH₂)_nNR₂]CO(CH₂)_nNR-aryl, N[(CH₂)_nNR₂]SO₂(CH₂)_n-aryl, N[(CH₂)_nXR]CO(CH₂)_n-heteroaryl, N[(CH₂)_nXR]CO(CH₂)_nX-heteroaryl, N[(CH₂)_nXR]SO₂(CH₂)_n-heteroaryl, N[(CH₂)_nNR₂]NRCOOR]CO(CH₂)_n-heteroaryl, N[(CH₂)_nNR₂]CO(CH₂)_nNR-heteroaryl, R¹ and R² together also denote -N-C(CF₃)=N-, -N-CR=N-, -N-N=N-,

Y denotes H, A, Hal

A denotes alkyl or cycloalkyl, in which one or more H atoms may be replaced by Hal,

Hal denotes F, Cl, Br or I

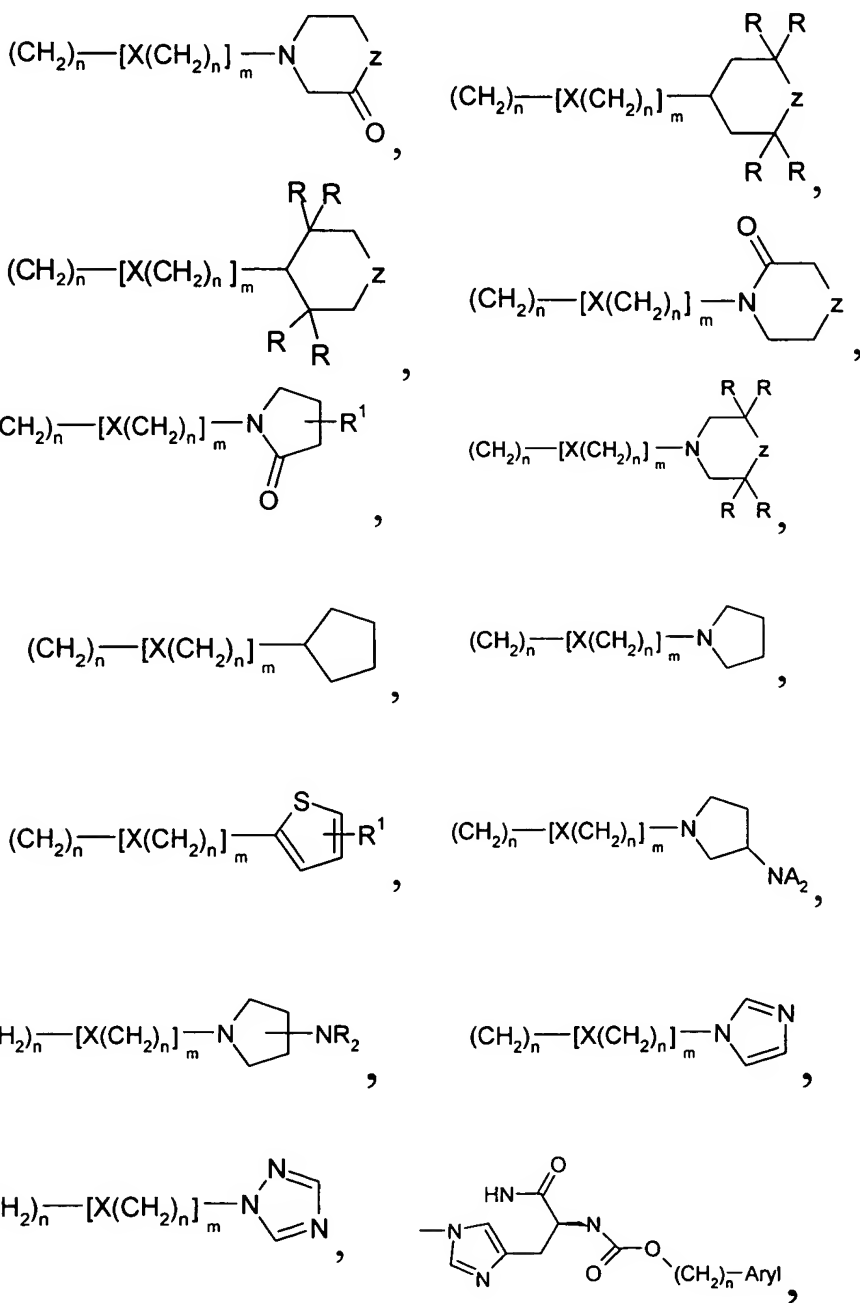
R denotes H or A, in the case of geminal radicals R together also -(CH₂)₅-, -(CH₂)₄- or -(CH₂)_n-X-(CH₂)_n, or -(CH₂)_n-Z-(CH₂)_n,

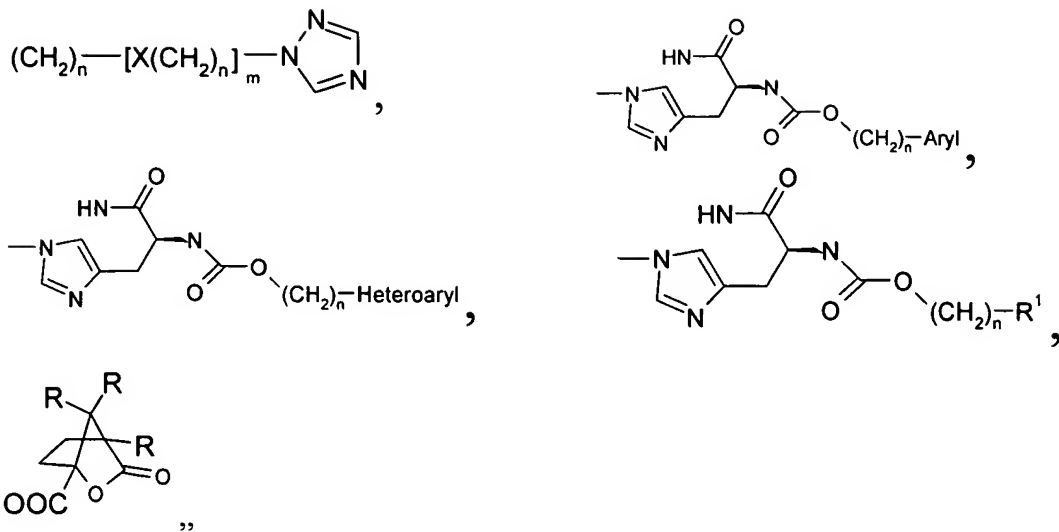
R⁴, R⁵, independently of one another, denote H or unsubstituted or mono- or poly-OR-, NO₂-, Hal-, CF₃-, OCF₃-, CN-, NR₂- or SR-, aryl- or heteroaryl-substituted N-pyrrolidone, -X-(CH₂)₂OR, -X-CO(CH₂)_nCH₃, -X-(CH₂)₂NR₂, R¹, S-aryl, O-aryl, CH₂Si(CH₃)₃, or together denote -X(CR₂)₂-, -X-(CR₂)₃-, -X-(CHCH₂OR)(CH₂)₂-, -X-(CHCH₂NR₂)(CH₂)₂-, -X(CH₂)₂NR₂, -(CR₂)₃-, -(CR₂)₄-, -CR=CR-CR=CR-, -XCHQ(CR₂)₂-, -XCHQCR₂-, R-N-(C=X)-N-R, -XC[(CH₂)_nOR]₂CH₂CH₂-,

X denotes O, S or NR,

Q denotes CH_2Hal , CHO , COR^a , CH_2R^a , CH_2OCOR^a ,
 CH_2NCOR^1 , $\text{CH}_2\text{N}(\text{R}^1)_2$, CH_2OR^1 , $\text{CH}_2\text{OCON}(\text{R}^1)_2$,
 $\text{CH}_2\text{OCOOR}^1$, $\text{CH}_2\text{NHCON}(\text{R}^1)_2$, $\text{CH}_2\text{NHCOOR}^1$,

R^a denotes





OR, NHR₂, NR₂, OR, NHR₂, NR₂, NR(CH₂)_n-aryl, NR(CH₂)_nOR,
 COOR, N-pyrrolidone radical, OCOR, NR(CH₂)_nNR₂,
 N[(CH₂)_nNR₂]CO(CH₂)_n-aryl, N[(CH₂)_nNHCOOR]CO-aryl, R¹,
 N[CH₂(CH₂)_nOR]₂, NR(CH₂)_nNCOOR, X(CH₂)_nX(CH₂)_nXR,
 NR(CH₂)_nX(CH₂)_nOH, NR(CH₂)_nO(CH₂)_nOH, (CH₂)_nCOOR,
 O(CO)NR(CH₂)_nOR, O(CO)(CH₂)_nNR₂, NR(CH₂)_nNR₂,
 N[(CH₂)_nNR₂]CO(CH₂)_n-aryl, N[(CH₂)_nXR]CO(CH₂)_n-aryl,
 N[(CH₂)_nXR]CO(CH₂)_n-heteroaryl, N[(CH₂)_nNR₂]CO(CH₂)_n-
 heteroaryl, N[(CH₂)_nNR₂]CO(CH₂)_nR¹, N(R)(CH₂)_nN(R)COOR,
 XCOO(CH₂)_nNR₂, OSO₂A, OSO₂CF₃, OSO₂Ar, OCONR₂,
 OCH₂(CH₂)_nNR₂

Z denotes CH₂, X, CHCONH₂, CH(CH₂)_nNRCOOR,
 CHNRCOOR, NCO, CH(CH₂)_nCOOR, NCOOR, CH(CH₂)_nOH,
 N(CH₂)_nOH, CHNH₂, CH(CH₂)_nNR₂, CH(CH₂)_nNR₂, C(OH)R,
 CHNCOR, CH(CH₂)_n-aryl, CH(CH₂)_n-heteroaryl, CH(CH₂)_nR¹,
 N(CH₂)_nCOOR, CH(CH₂)_nX(CH₂)_n-aryl, CH(CH₂)_nX(CH₂)_n-
 heteroaryl, N(CH₂)_nCONR₂, XCONR(CH₂)_nNR₂,
 N[(CH₂)_nXCOOR]CO(CH₂)_n-aryl, N[(CH₂)_nXR]CO(CH₂)_n-aryl,
 N[(CH₂)_nXR]CO(CH₂)_nX-aryl, N[(CH₂)_nXR]SO₂(CH₂)_n-aryl,

N[(CH₂)_nNRCOOR]CO(CH₂)_n-aryl, N[(CH₂)_nNR₂]CO(CH₂)_n-aryl, N[(CH₂)_nNR₂]CO(CH₂)_nNR-aryl, N[(CH₂)_nNR₂]SO₂(CH₂)_n-aryl, N[(CH₂)_nXR]CO(CH₂)_n-heteroaryl, N[(CH₂)_nXR]CO(CH₂)_nX-heteroaryl, N[(CH₂)_nXR]SO₂(CH₂)_n-heteroaryl, N[(CH₂)_nNRCOOR]CO(CH₂)_n-heteroaryl, N[(CH₂)_nNR₂]CO(CH₂)_n-heteroaryl, N[(CH₂)_nNR₂]SO₂(CH₂)_n-heteroaryl, O(CH₂)_nNR₂, X(CH₂)_nNR₂, NCO(CH₂)_nNR₂,

R⁶ denotes aryl or heteroaryl, each of which is unsubstituted or mono- or polysubstituted by aryl or heteroaryl, each of which may be substituted by Hal, NO₂, CN, A, OR, OCOR, COR, NR₂, CF₃, OCF₃, OCH(CF₃)₂, or by Hal, NO₂, CN, OR, A, -(CY₂)_n-OR, -OCOR, -(CY₂)_n-CO₂R, -(CY₂)_n-CN, -NCOR, -COR or -(CY₂)_n-NR₂,

R⁷ denotes (C=O)-R, (C=O)-NR₂, (C=O)-OR, H or A

m denotes 0, 1 or 2

and

n denotes 0, 1, 2, 3, 4, 5, 6 or 7,

and pharmaceutically usable derivatives, solvates, tautomers, salts and stereoisomers thereof, including mixtures thereof in all ratios.

The invention also relates to the optically active forms, the enantiomers, the racemates, the diastereomers and the hydrates and solvates of these compounds. The term solvates of the compounds is taken to mean adductions of inert solvent molecules onto the compounds of the formula I which form owing to their mutual attractive force. Solvates are, for example, mono- or dihydrates or alkoxides.

The term pharmaceutically usable derivatives is taken to mean, for example, the salts of the compounds according to the invention and also so-called prodrug compounds.

The term prodrug derivatives is taken to mean compounds of the formula I which have been modified by means of, for example, alkyl or acyl groups, sugars or oligopeptides and which are rapidly cleaved in the organism to form the effective compounds according to the invention.

These also include biodegradable polymer derivatives of the compounds according to the invention, as described, for example, in Int. J. Pharm. 115, 61-67 (1995).

Similar compounds are described, for example, in Tetrahedron Lett. **1988**, 29, 5855-5858, Tetrahedron Lett. **2003**, 44, 217-219, J. Org. Chem. **1997**, 62, 4880-4882, J. Org. Chem. **1999**, 64, 6462-6467, Chem. Lett. **1995**, 423-424, J. Org. Chem. **2000**, 65, 5009-5013, Chem. Lett. **2003**, 32, 222-223, US2003149069A1, but are not mentioned in connection with cancer treatments and/or do not contain the features that are essential to the invention.

The expression "effective amount" denotes the amount of a medicament or of a pharmaceutical active ingredient which causes in a tissue, system, animal or human a biological or medical response which is sought or desired, for example, by a researcher or physician.

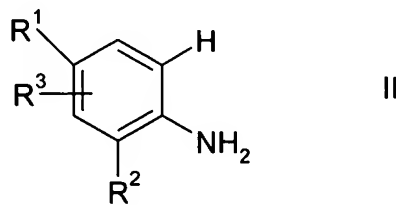
In addition, the expression "therapeutically effective amount" denotes an amount which causes at least one of the following effects in a human or mammal (compared with a subject who has not received this amount): improvement in the healing treatment, healing, prevention or elimination of a disease, syndrome, condition, complaint, disorder or side-effects or also the reduction in the progress of a disease, condition or disorder.

The expression "therapeutically effective amount" also encompasses the amounts which are effective for increasing or strengthening normal physiological function.

The invention also relates to the use of mixtures of the compounds according to the invention, for example mixtures of two diastereomers, for example in the ratio 1:1, 1:2, 1:3, 1:4, 1:5, 1:10, 1:100 or 1:1000.

These are particularly preferably mixtures of stereoisomeric compounds.

The invention relates to the compounds of the formula I and salts thereof and to a process for the preparation of compounds of the formula I according to the patent claims and pharmaceutically usable derivatives, salts, solvates and stereoisomers thereof, characterised in that a compound of the formula II



in which R¹, R² and R³ have the meanings indicated above,

is reacted with a compound of the formula III

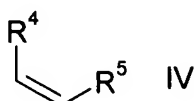


in which

R⁶ has the meaning indicated above,

and

with a compound of the formula IV, the double-bond isomer thereof (E isomer) or mixtures thereof



in which R^4 and R^5 have the meanings indicated above, preferably in the presence of a protonic acid or Lewis acid, such as, for example, trifluoroacetic acid, hexafluoroisopropanol, bismuth(III) chloride, ytterbium(III) triflate, scandium(III) triflate or cerium(IV) ammonium nitrate,

and a radical other than H is optionally introduced by conventional methods for R^7 .

The mixtures of diastereomers and enantiomers of the compounds of the formula I which may be obtained by the process described above are preferably separated by chromatography or crystallisation.

If desired, the bases and acids of the formula I obtained by the process described above are converted into their salts.

Above and below, the radicals R, R^1 , R^2 , R^3 , R^4 , R^5 , R^6 , R^7 , X, Y, Q, R^a , Z, W, m and n have the meanings indicated for the formula I, unless expressly indicated otherwise. If individual radicals occur a number of times within a compound, the radicals adopt the meanings indicated, independently of one another.

A denotes alkyl, is preferably unbranched (linear) or branched, and has 1, 2, 3, 4, 5, 6, 7, 8, 9 or 10 C atoms. A preferably denotes methyl, further-

more ethyl, propyl, isopropyl, butyl, isobutyl, sec-butyl or tert-butyl, furthermore also pentyl, 1-, 2- or 3-methylbutyl, 1,1-, 1,2- or 2,2-dimethylpropyl, 1-ethylpropyl, hexyl, 1-, 2-, 3- or 4-methylpentyl, 1,1-, 1,2-, 1,3-, 2,2-, 2,3- or 3,3-dimethylbutyl, 1- or 2-ethylbutyl, 1-ethyl-1-methylpropyl, 1-ethyl-2-methylpropyl, 1,1,2- or 1,2,2-trimethylpropyl, furthermore preferably, for example, trifluoromethyl.

A very particularly preferably denotes alkyl having 1, 2, 3, 4, 5 or 6 C atoms, preferably methyl, ethyl, propyl, isopropyl, butyl, isobutyl, sec-butyl, tert-butyl, pentyl, hexyl, trifluoromethyl, pentafluoroethyl or 1,1,1-trifluoroethyl. A also denotes cycloalkyl.

Cycloalkyl preferably denotes cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl or cycloheptyl, but in particular cyclopentyl.

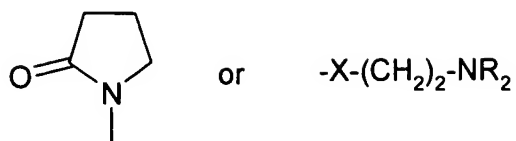
R¹ preferably denotes A, CF₃, OCF₃, SA, SCN, CH₂CN, -OCOA, Hal, SCF₃, preferably also t-butyl, -CH(CH₃)CH₂CH₃, isopropyl, ethyl or methyl. In particular, R¹ denotes t-butyl, isopropyl, ethyl, CF₃, methyl, Br, Cl, SCF₃, CH(CH₃)CH₂CH₃, n-propyl, OCH₃, SCH₃, n-butyl, -SCN, CH₂CN. R¹ particularly preferably denotes t-butyl, isopropyl, ethyl or CF₃.

R² preferably denotes Hal, A or OA, in particular Br, cyclopropyl, OCH₃. Particular preference is furthermore given to H or F.

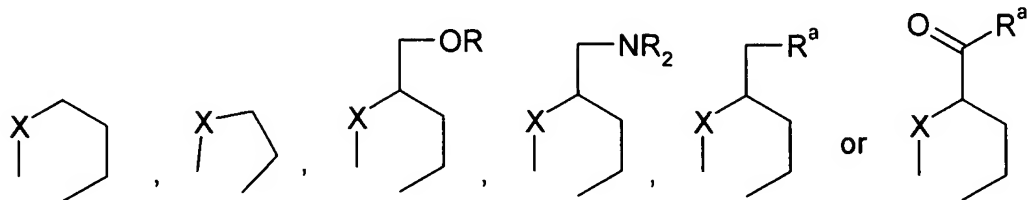
R³ preferably denotes H or A, in particular H. R³ is preferably in the 5-position. In particular, R³ denotes H or F.

In particularly preferred compounds of the formula I, R² and R³ simultaneously have the meaning H. In further preferred compounds of the formula I, one of the radicals R² and R³ has the meaning H and the other radical has the meaning F.

R⁴ preferably denotes one of the following groups if R⁵ denotes H:

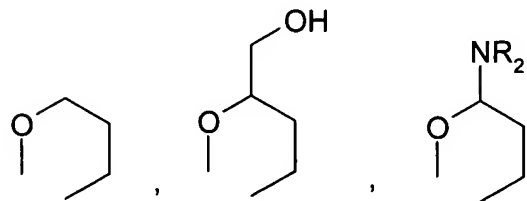


R⁵ preferably denotes H or, together with R⁴, adopts one of the following meanings:



in which X, R and R^a have the meaning indicated above.

R⁴, together with R⁵, particularly preferably adopts one of the following meanings:

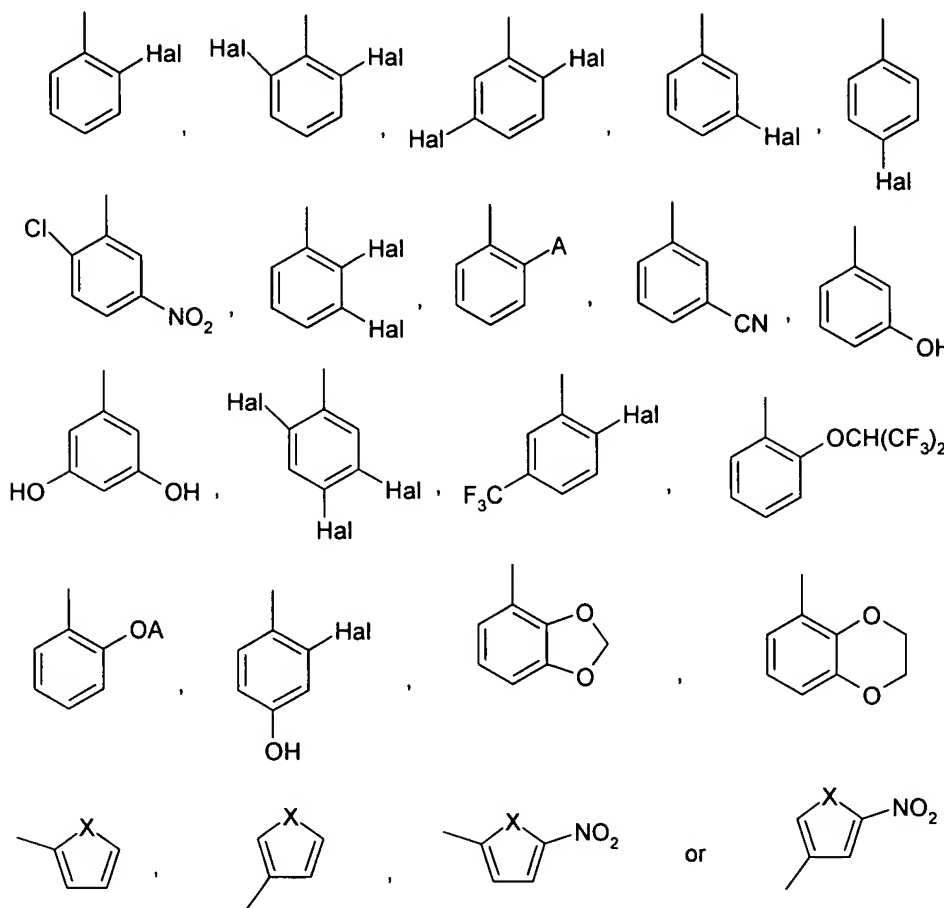


in which R has the meaning indicated above.

R^a preferably denotes 1-piperazinyl, N-morpholinyl, NHR or NR₂.

If the radicals and indices, such as, for example, n, occur more than once, the radicals and indices may, independently of one another, adopt different values. In the radicals R^a, n preferably denotes 2 or 0 and m preferably denotes 0.

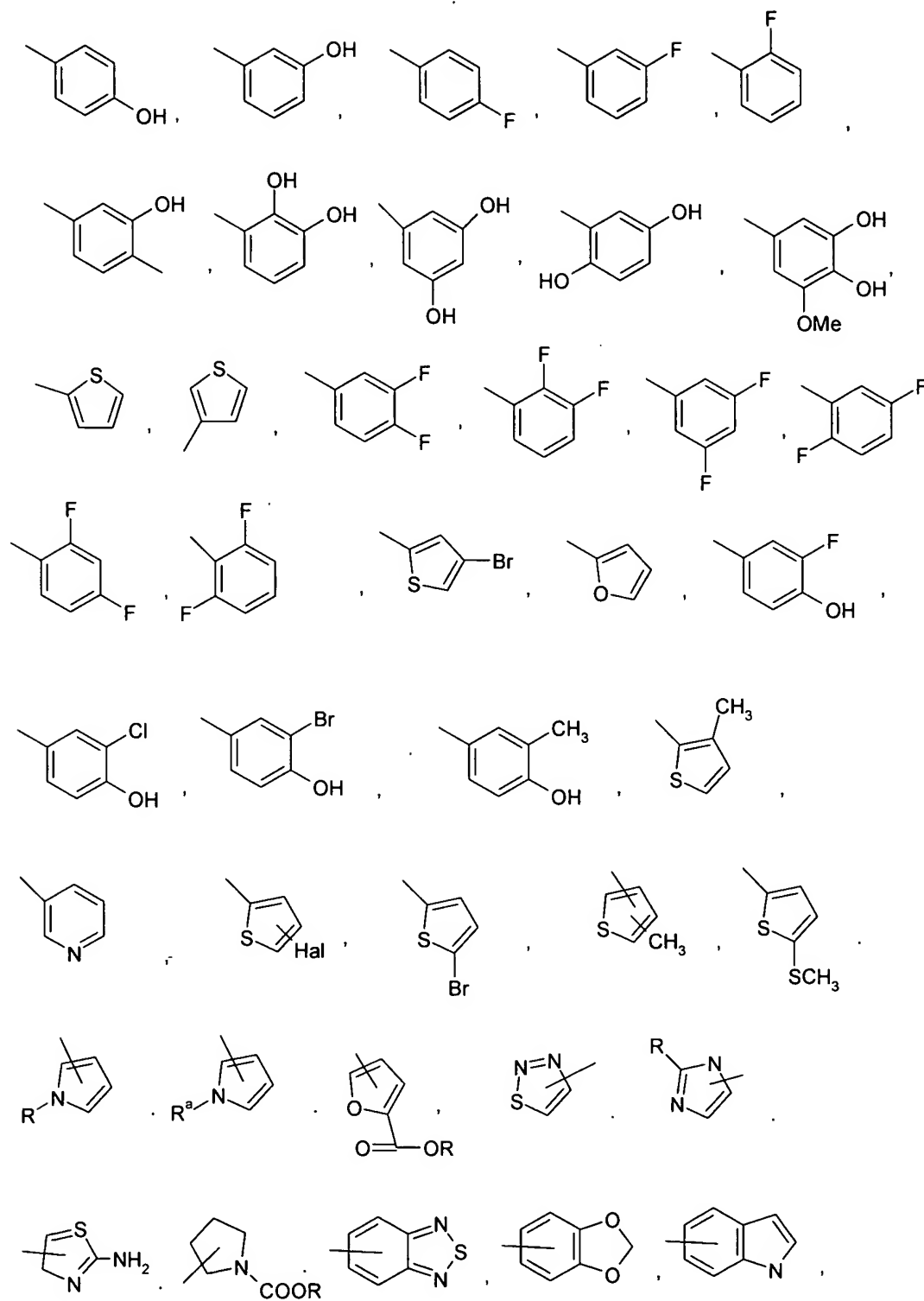
R⁶ preferably denotes phenyl, 2-, 3- or 4-pyridyl, pyrimidyl, furyl or thienyl, each of which is unsubstituted or mono- or polysubstituted by Hal, CN, NO₂, OH, CF₃, OCH(CF₃)₂, OCOCH₃ or A. R⁶ is preferably not a heteroaromatic radical. In particular, R⁶ denotes one of the following groups:



in which

X denotes O, S or NR and in particular O or S, A has the meaning indicated above, but preferably denotes methyl, and Hal preferably denotes F or Cl.

Particular preference is furthermore given to compounds of the formula I in which R⁶ has one of the following meanings:



R^7 preferably denotes H or A, in particular H.

Aryl preferably denotes phenyl, naphthyl or biphenyl, each of which is unsubstituted or mono-, di- or trisubstituted by Hal, A, OH, OA, NH₂, NO₂, CN, COOH, COOA, CONH₂, NHCOA, NHCONH₂, NSO₂A, CHO, COA, SO₂NH₂, SO₂A, -CH₂-COOH or -OCH₂-COOH.

Aryl preferably denotes phenyl, o-, m- or p-tolyl, o-, m- or p-ethylphenyl, o-, m- or p-propylphenyl, o-, m- or p-isopropylphenyl, o-, m- or p-tert-butylphenyl, o-, m- or p-hydroxyphenyl, o-, m- or p-methoxyphenyl, o-, m- or p-nitrophenyl, o-, m- or p-aminophenyl, o-, m- or p-(N-methylamino)phenyl, o-, m- or p-(N-methylaminocarbonyl)phenyl, o-, m- or p-acetamidophenyl, o-, m- or p-methoxyphenyl, o-, m- or p-ethoxyphenyl, o-, m- or p-ethoxy-carbonylphenyl, o-, m- or p-(N,N-dimethylamino)phenyl, o-, m- or p-(N,N-dimethylaminocarbonyl)phenyl, o-, m- or p-(N-ethylamino)phenyl, o-, m- or p-(N,N-diethylamino)phenyl, o-, m- or p-fluorophenyl, o-, m- or p-bromo-phenyl, o-, m- or p-chlorophenyl, o-, m- or p-(methylsulfonamido)phenyl, o-, m- or p-(methylsulfonyl)phenyl, furthermore preferably 2,3-, 2,4-, 2,5-, 2,6-, 3,4- or 3,5-difluorophenyl, 2,3-, 2,4-, 2,5-, 2,6-, 3,4- or 3,5-dichlorophenyl, 2,3-, 2,4-, 2,5-, 2,6-, 3,4- or 3,5-dibromophenyl, 2,4- or 2,5-dinitrophenyl, 2,5- or 3,4-dimethoxyphenyl, 3-nitro-4-chlorophenyl, 3-amino-4-chloro-, 2-amino-3-chloro-, 2-amino-4-chloro-, 2-amino-5-chloro- or 2-amino-6-chlorophenyl, 2-nitro-4-N,N-dimethylamino- or 3-nitro-4-N,N-dimethylaminophenyl, 2,3-diaminophenyl, 2,3,4-, 2,3,5-, 2,3,6-, 2,4,6- or 3,4,5-trichlorophenyl, 2,4,6-trimethoxyphenyl, 2-hydroxy-3,5-dichlorophenyl, p-iodophenyl, 3,6-dichloro-4-aminophenyl, 4-fluoro-3-chlorophenyl, 2-fluoro-4-bromophenyl, 2,5-difluoro-4-bromophenyl, 3-bromo-6-methoxyphenyl, 3-chloro-6-methoxyphenyl, 3-chloro-4-acetamidophenyl, 3-fluoro-4-methoxyphenyl, 3-amino-6-methylphenyl, 3-chloro-4-acetamidophenyl or 2,5-dimethyl-4-chlorophenyl.

Heteroaryl preferably denotes a mono- or bicyclic aromatic heterocycle having one or more N, O and/or S atoms which is unsubstituted or mono-, di- or trisubstituted by Hal, A, NO₂, NHA, NA₂, OA, COOA or CN.

Heteroaryl particularly preferably denotes a monocyclic saturated or aromatic heterocycle having one N, S or O atom, which may be unsubstituted or mono-, di- or trisubstituted by Hal, A, NHA, NA₂, NO₂, COOA or benzyl.

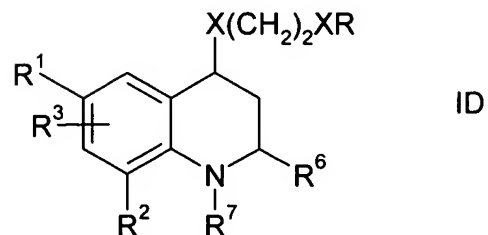
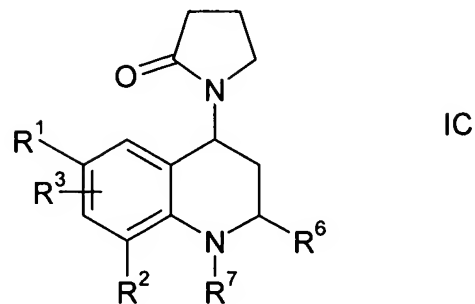
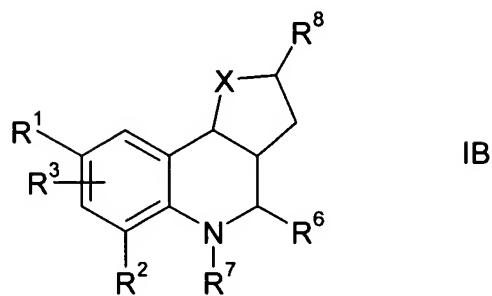
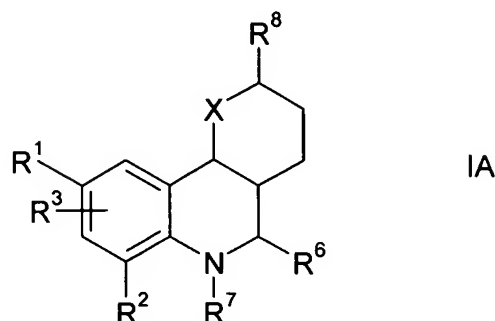
Irrespective of further substitutions, unsubstituted heteroaryl denotes, for example, 2- or 3-furyl, 2- or 3-thienyl, 1-, 2- or 3-pyrrolyl, 1-, 2-, 4- or 5-imidazolyl, 1-, 3-, 4- or 5-pyrazolyl, 2-, 4- or 5-oxazolyl, 3-, 4- or 5-isoxazolyl, 2-, 4- or 5-thiazolyl, 3-, 4- or 5-isothiazolyl, 2-, 3- or 4-pyridyl, 2-, 4-, 5- or 6-pyrimidinyl, furthermore preferably 1,2,3-triazol-1-, -4- or -5-yl, 1,2,4-triazol-1-, -3- or 5-yl, 1- or 5-tetrazolyl, 1,2,3-oxadiazol-4- or -5-yl, 1,2,4-oxadiazol-3- or -5-yl, 1,3,4-thiadiazol-2- or -5-yl, 1,2,4-thiadiazol-3- or -5-yl, 1,2,3-thiadiazol-4- or -5-yl, 3- or 4-pyridazinyl, pyrazinyl, 1-, 2-, 3-, 4-, 5-, 6- or 7-indolyl, 4- or 5-isoindolyl, 1-, 2-, 4- or 5-benzimidazolyl, 1-, 3-, 4-, 5-, 6- or 7-benzopyrazolyl, 2-, 4-, 5-, 6- or 7-benzoxazolyl, 3-, 4-, 5-, 6- or 7-benzisoxazolyl, 2-, 4-, 5-, 6- or 7-benzothiazolyl, 2-, 4-, 5-, 6- or 7-benzisothiazolyl, 4-, 5-, 6- or 7-benz-2,1,3-oxadiazolyl, 2-, 3-, 4-, 5-, 6-, 7- or 8-quinolyl, 1-, 3-, 4-, 5-, 6-, 7- or 8-isoquinolyl, 3-, 4-, 5-, 6-, 7- or 8-cinnolinyl, 2-, 4-, 5-, 6-, 7- or 8-quinazolinyl, 5- or 6-quinoxalinyl, 2-, 3-, 5-, 6-, 7- or 8-2H-benzo-1,4-oxazinyl, furthermore preferably 1,3-benzodioxol-5-yl, 1,4-benzodioxan-6-yl, 2,1,3-benzothiadiazol-4- or -5-yl or 2,1,3-benzoxadiazol-5-yl.

Hal preferably denotes F, Cl or Br, but also I, particularly preferably F or Cl.

Throughout the invention, all radicals which occur more than once may be identical or different, i.e. are independent of one another.

The compounds of the formula I may have one or more chiral centres and therefore exist in various stereoisomeric forms. The formula I encompasses all these forms.

Particularly preferred compounds of the formula I are those of the sub-formulae IA to ID:



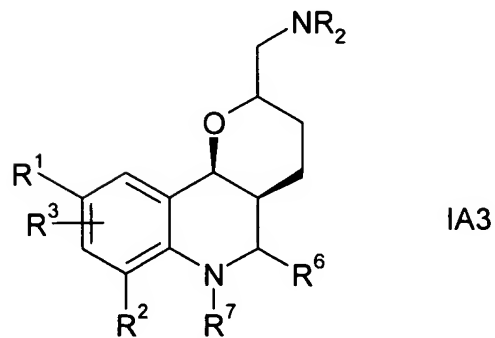
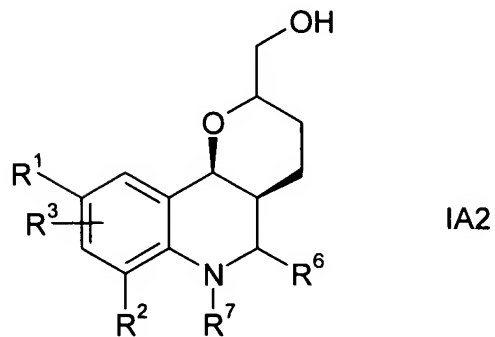
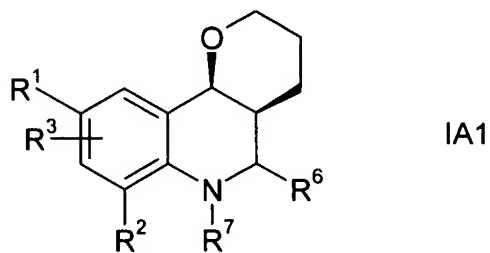
in which

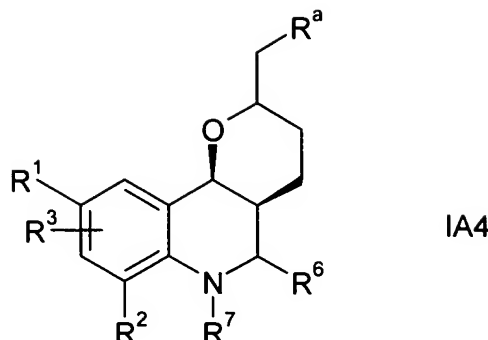
R, R¹, R², R⁶, R⁷ and X have the meanings indicated above

and

R^8 denotes H, CH_2OR , CH_2NR_2 , CH_2R^a , COR^a .

Particularly preferred compounds of the formula IA are those of the sub-formulae IA1 to IA4:



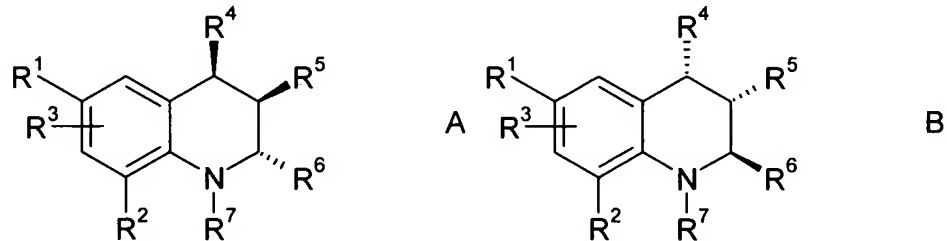


in which R , R^a , R^1 , R^2 , R^3 , R^6 and R^7 have the meanings indicated above.

In particularly preferred compounds of the formula IB, R⁸ has the meaning H.

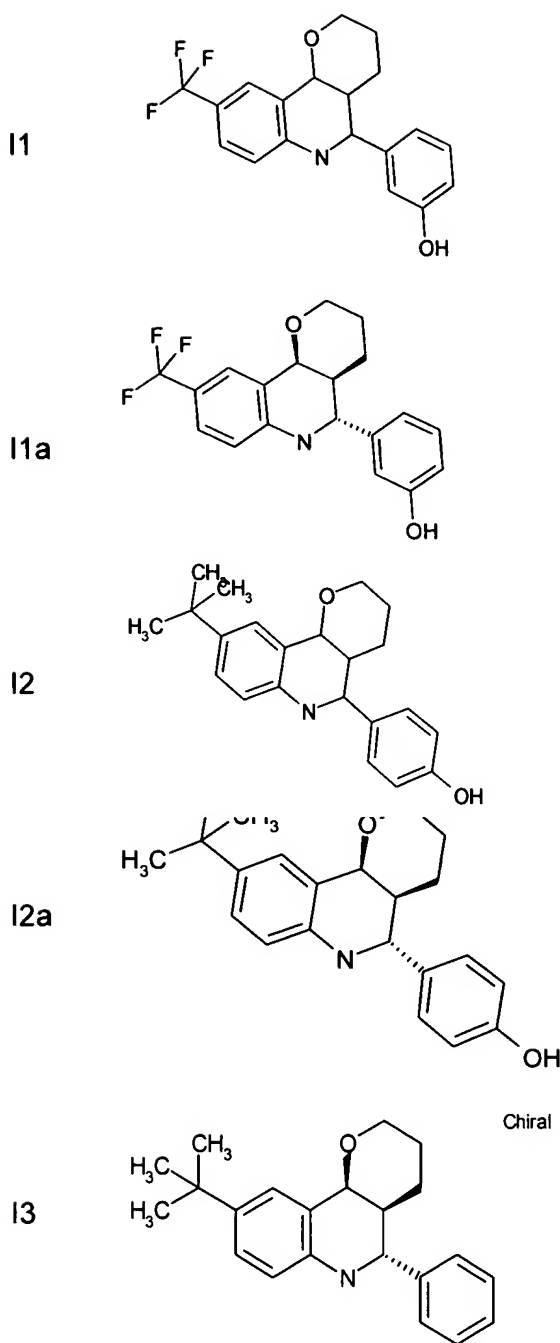
The radicals R⁴ and R⁵ are particularly preferably in the cis-position to one another. The radical R⁶ is furthermore preferably in the trans-position to the radical R⁵.

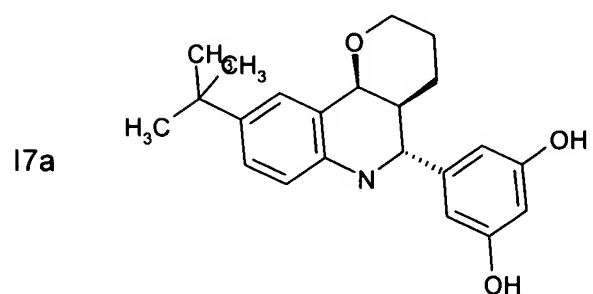
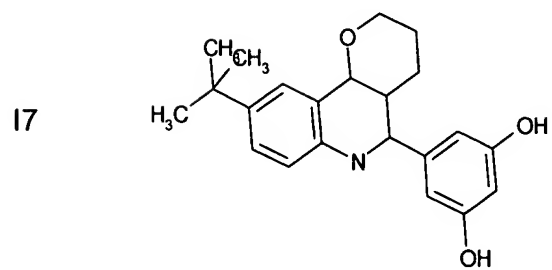
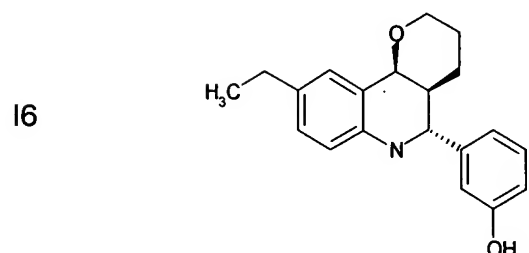
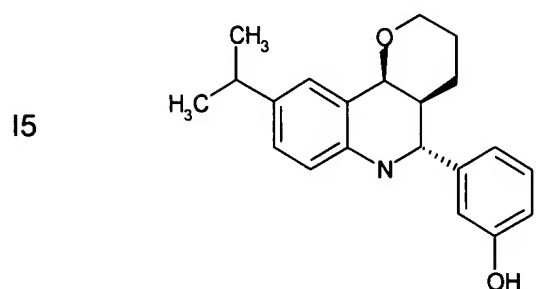
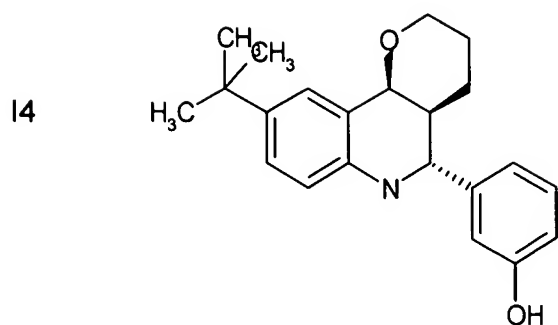
Preference is given here to a compound of the formula A or B having the following structure:

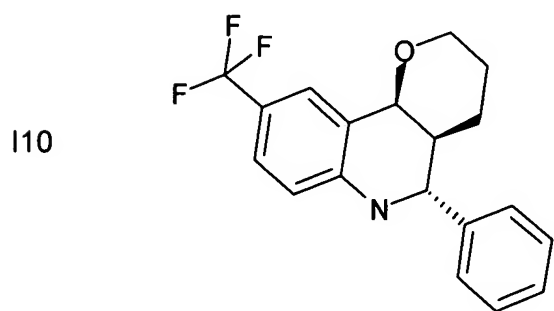
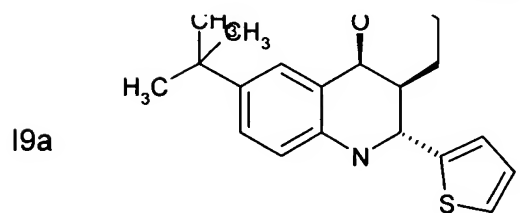
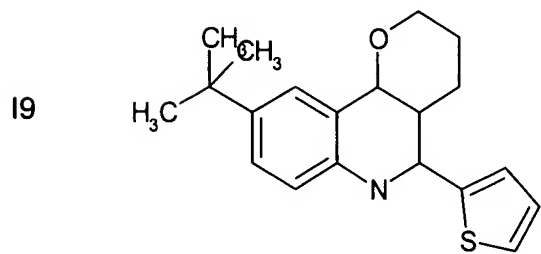
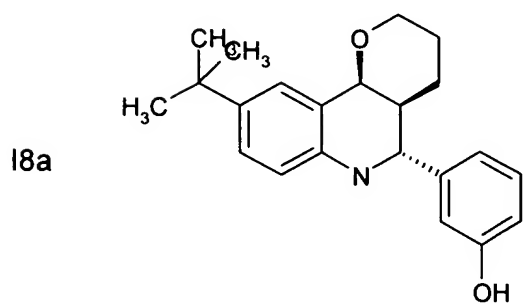
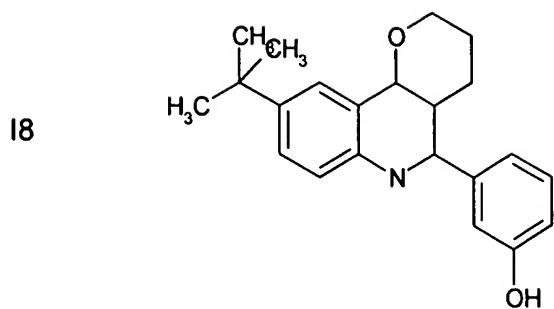


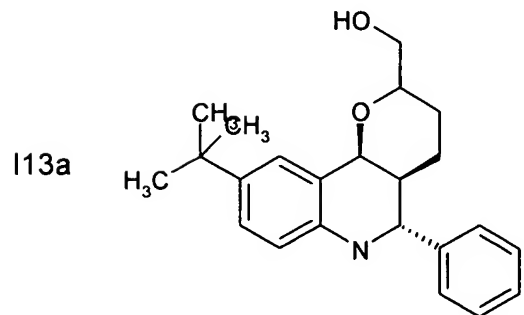
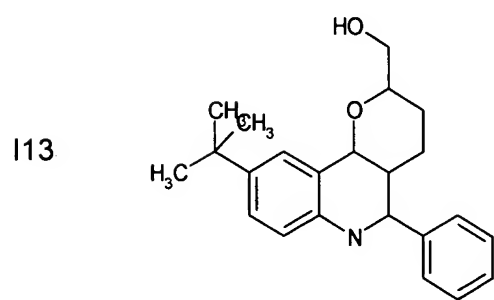
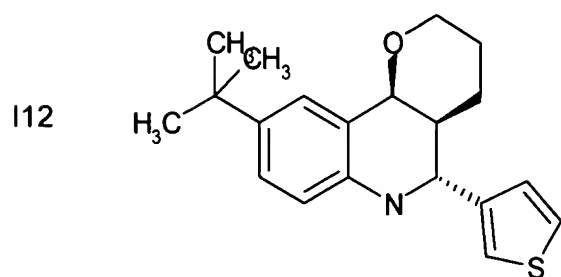
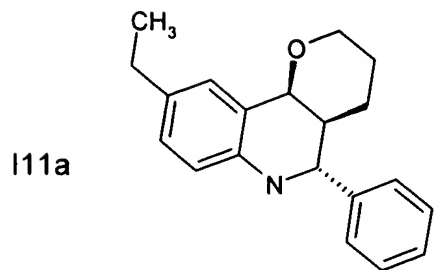
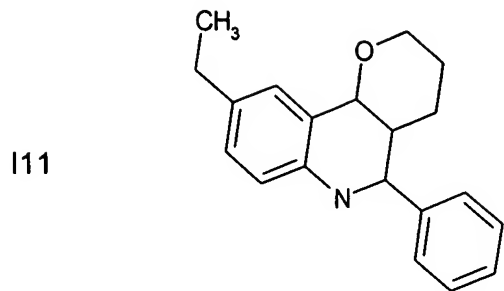
and the racemate thereof or other mixtures of the enantiomers.

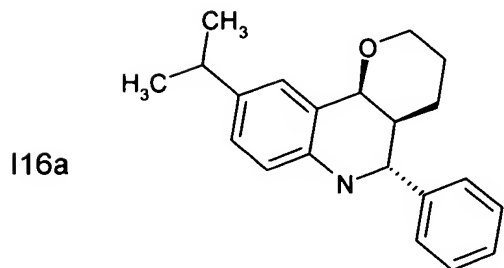
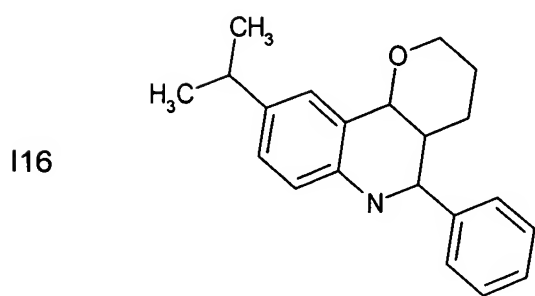
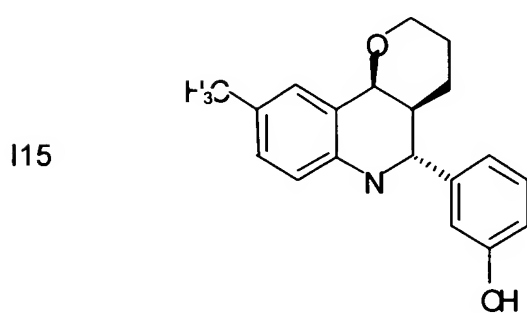
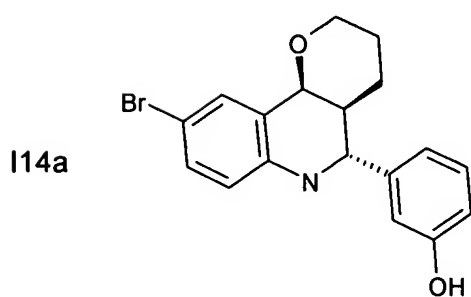
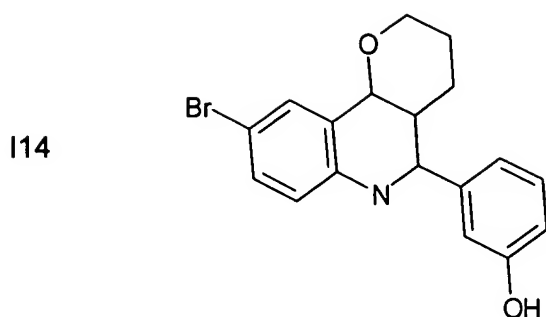
Accordingly, the invention relates, in particular, to the compounds of the formula I in which at least one of the said radicals has one of the preferred meanings indicated above. Some preferred groups of compounds may be expressed by the following sub-formulae I1 to I45a:

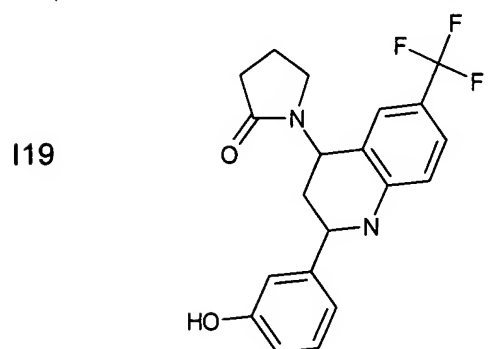
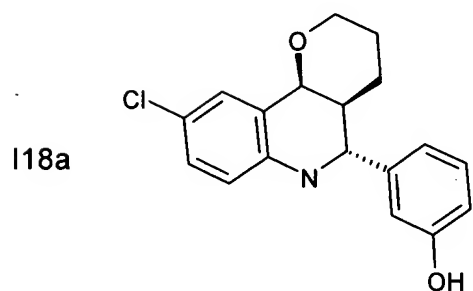
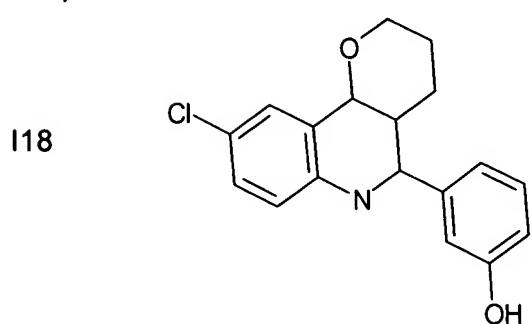
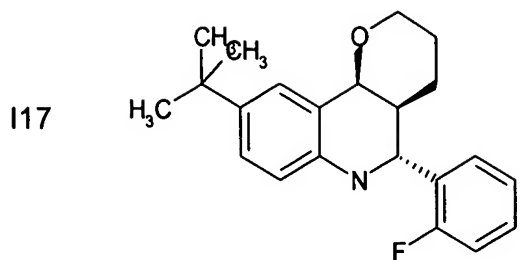


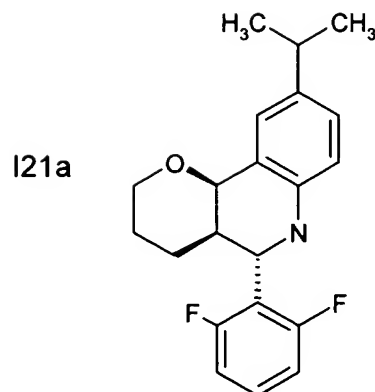
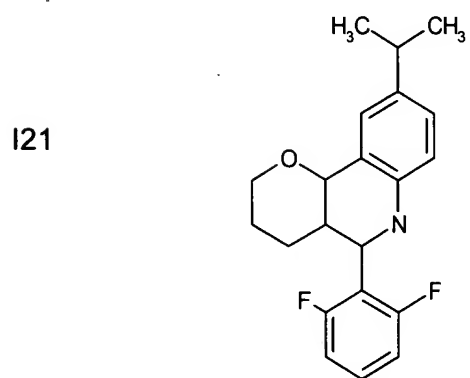
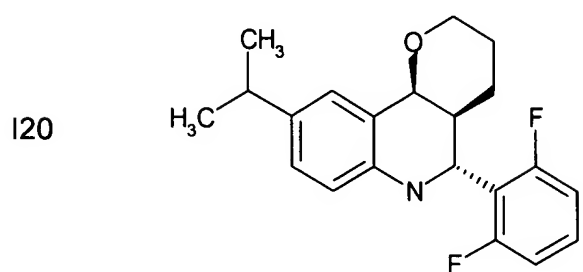
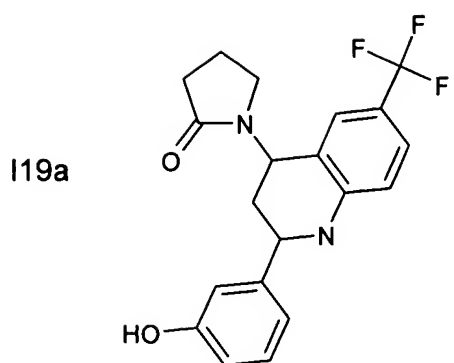


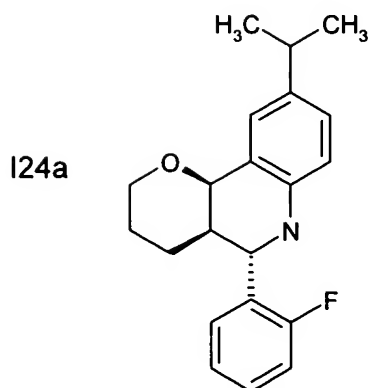
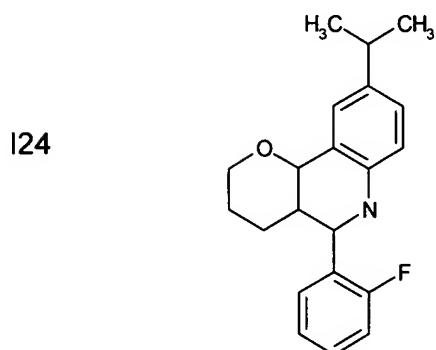
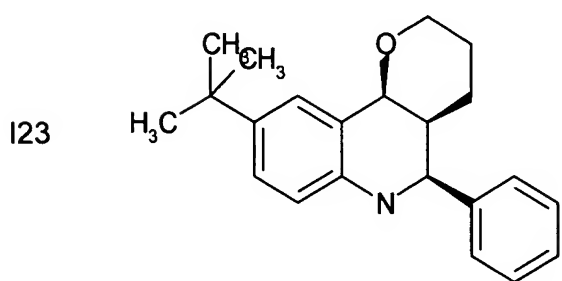
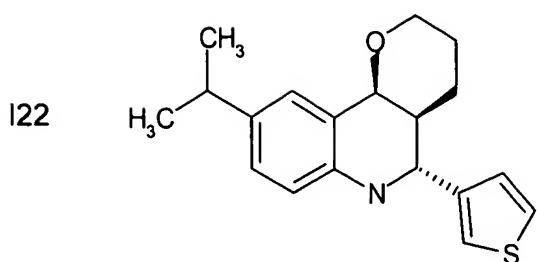


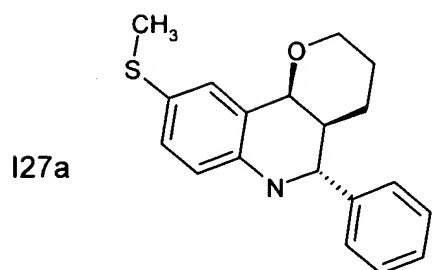
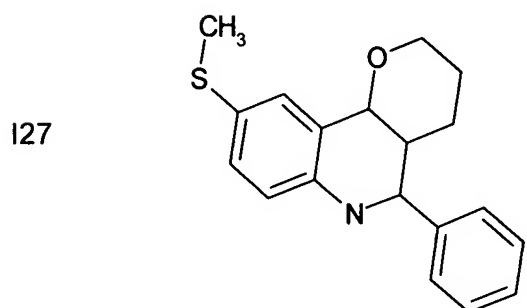
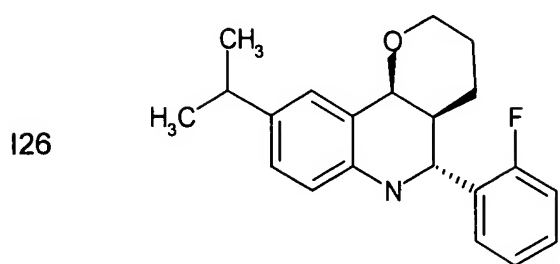
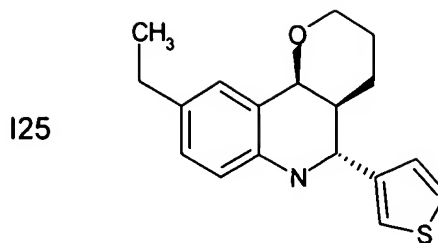


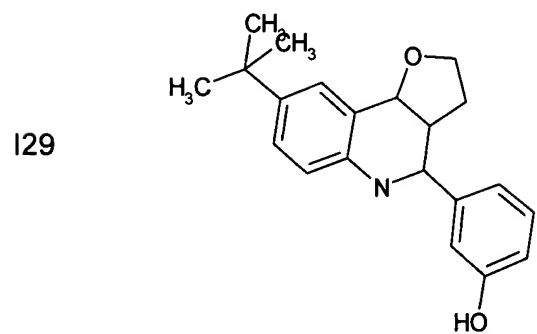
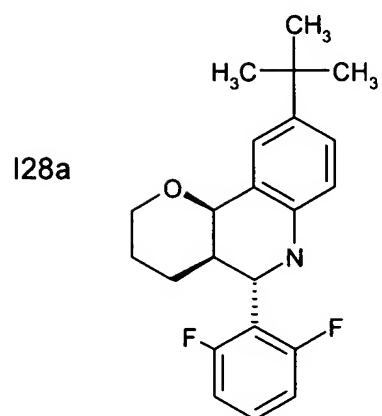
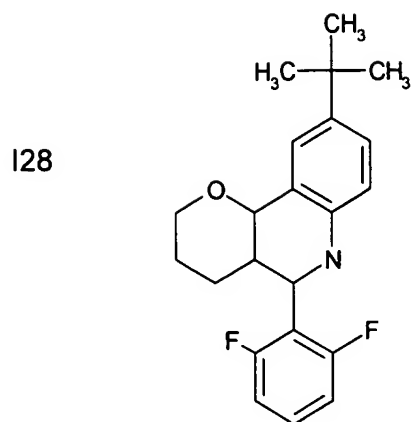


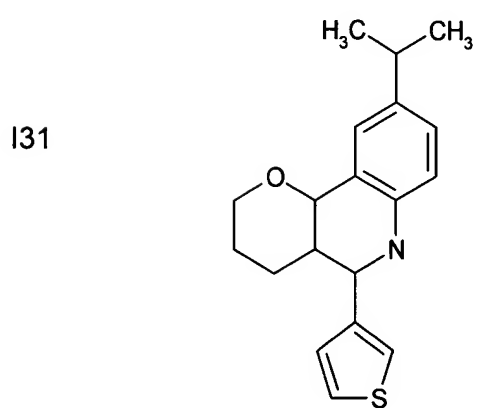
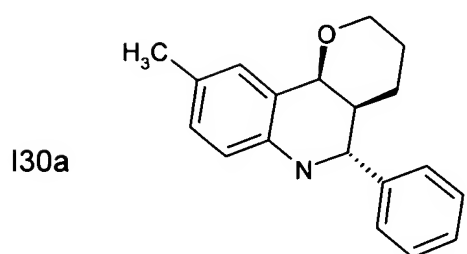
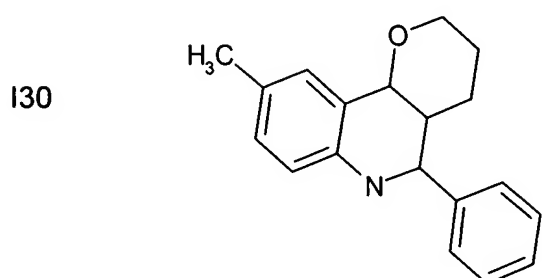
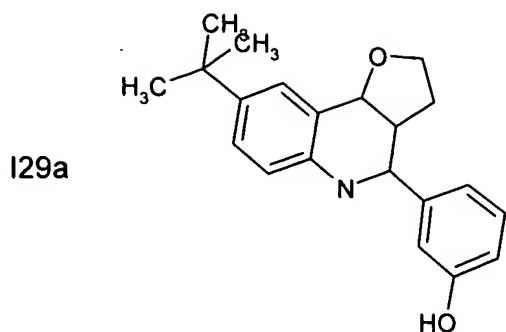


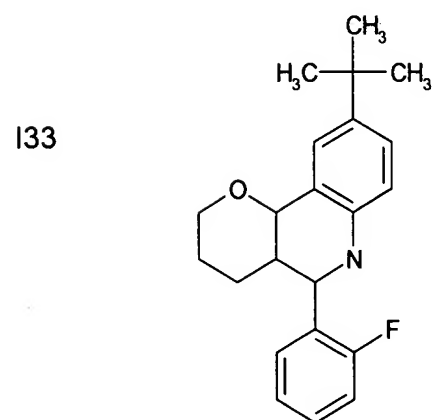
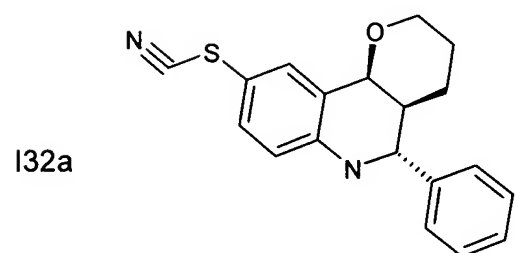
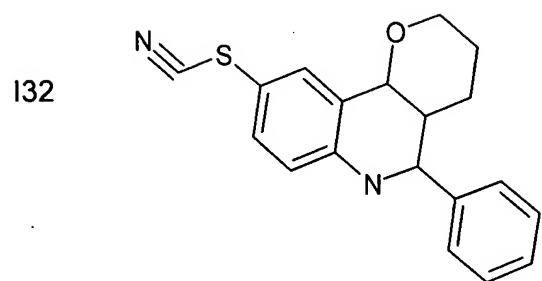
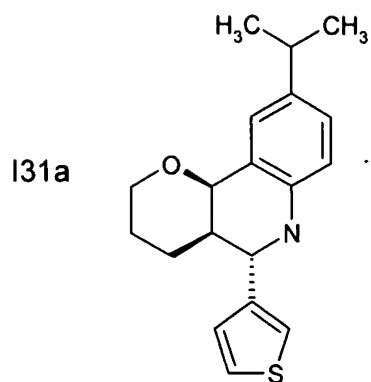


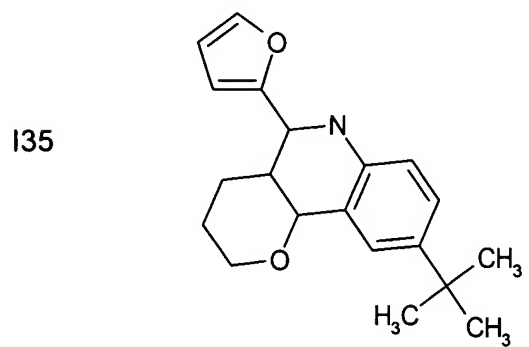
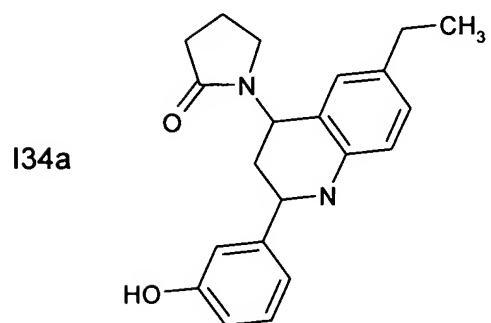
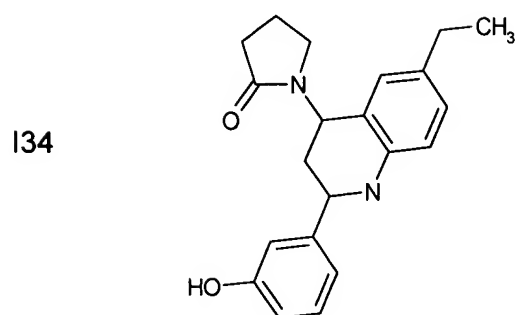
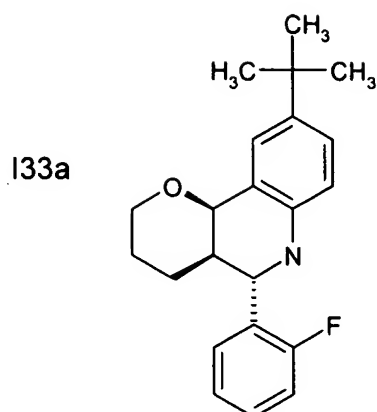


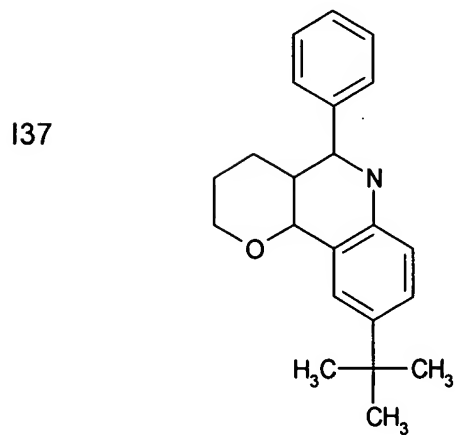
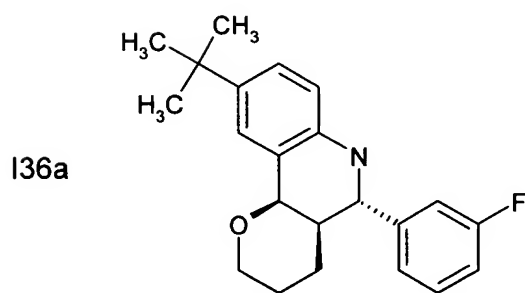
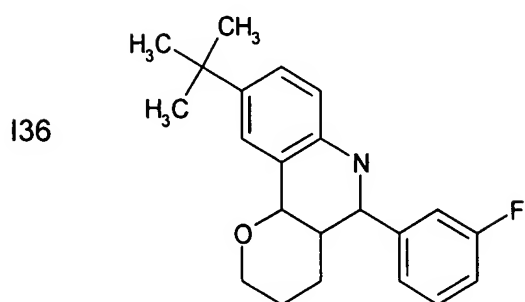
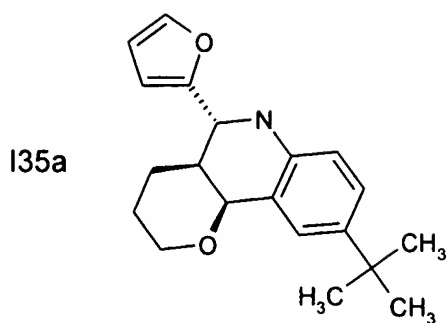


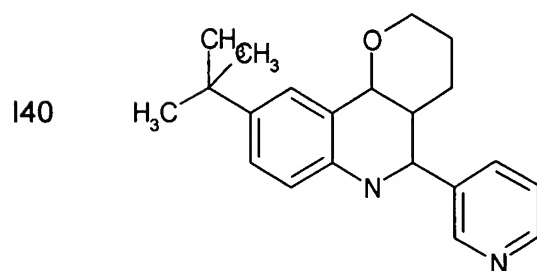
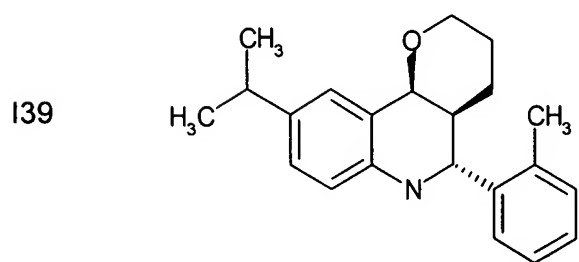
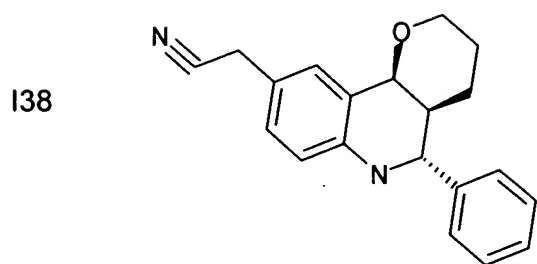
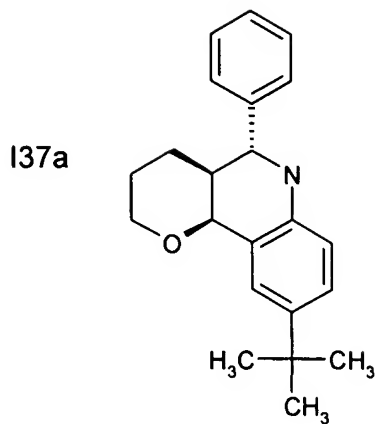


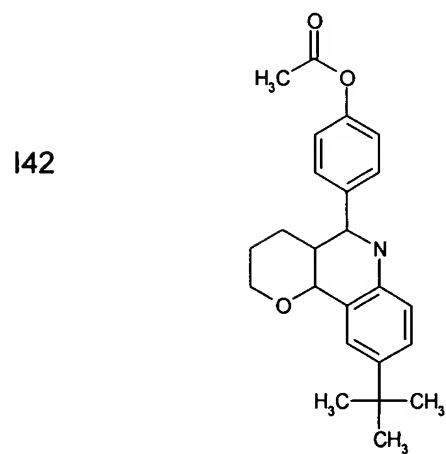
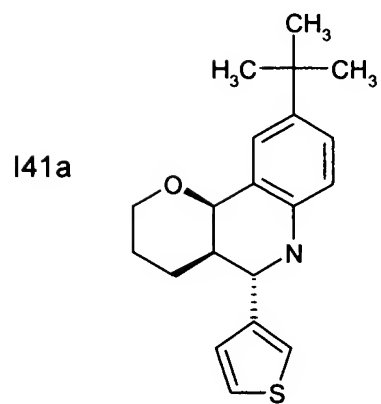
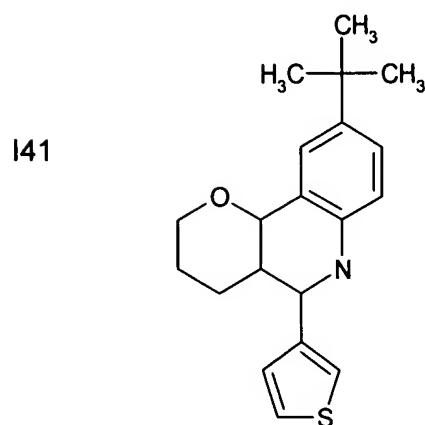
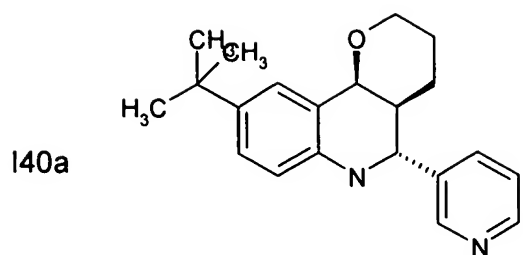


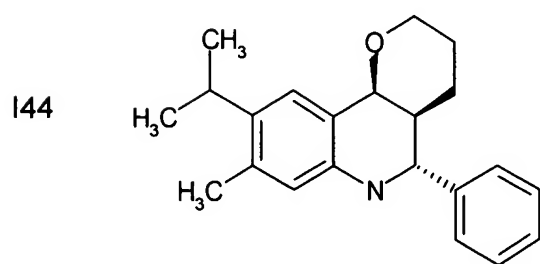
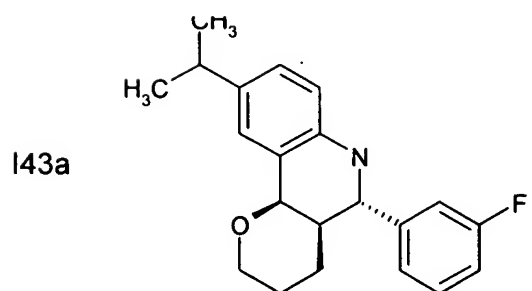
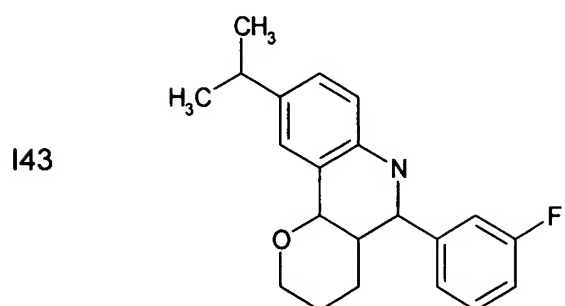
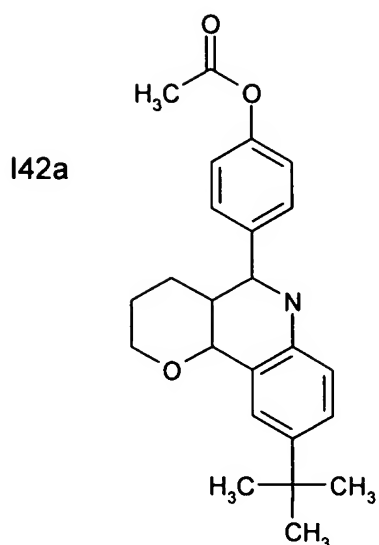


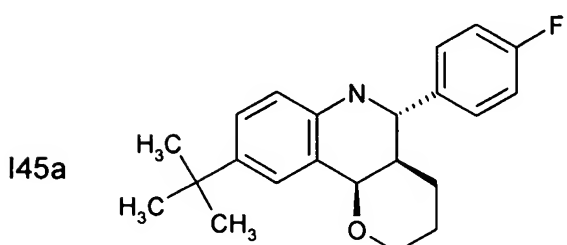
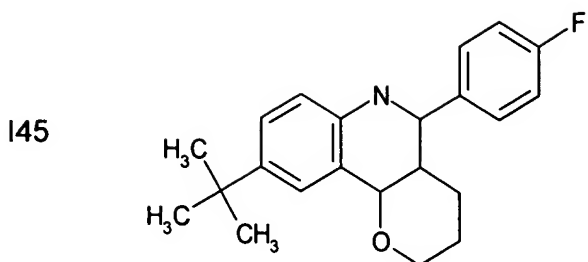












The compounds of the formula I and also the starting materials for their preparation are, in addition, prepared by methods known per se, as described in the literature (for example in the standard works, such as Houben-Weyl, Methoden der organischen Chemie [Methods of Organic Chemistry], Georg-Thieme-Verlag, Stuttgart), to be precise under reaction conditions which are known and suitable for the said reactions. Use may also be made here of variants known per se which are not mentioned here in greater detail.

If desired, the starting materials may also be formed in situ so that they are not isolated from the reaction mixture, but instead are immediately converted further into the compounds of the formula I.

The reaction is generally carried out in an inert solvent, preferably in the presence of a protonic acid or Lewis acid, such as TFA, HFIP, bismuth(III) salts, ytterbium(III) salts or CAN. Depending on the conditions used, the reaction time is between a few minutes and 14 days, the reaction tempera-

ture is between about 0° and 180°, normally between 0° and 100°, particularly preferably between 15° and 35°C.

Suitable inert solvents are, for example, hydrocarbons, such as hexane, petroleum ether, benzene, toluene or xylene; chlorinated hydrocarbons, such as trichloroethylene, 1,2-dichloroethane, carbon tetrachloride, chloroform or dichloromethane; nitriles, such as acetonitrile; carbon disulfide; carboxylic acids, such as formic acid or acetic acid; nitro compounds, such as nitromethane or nitrobenzene, or mixtures of the said solvents.

Compounds of the formula I in which R⁷ has a meaning other than H are preferably prepared by alkylation or acylation from the compounds of the formula I in which R⁷ denotes H.

If desired, a functionally modified amino and/or hydroxyl group in a compound of the formula I can be liberated by solvolysis or hydrogenolysis by conventional methods. This can be carried out, for example, using NaOH or KOH in water, water/THF or water/dioxane at temperatures between 0 and 100°.

The reduction of an ester to the aldehyde or alcohol or the reduction of a nitrile to the aldehyde or amine is carried out by methods as are known to the person skilled in the art and are described in standard works of organic chemistry.

The said compounds according to the invention can be used in their final non-salt form. On the other hand, the present invention also relates to the use of these compounds in the form of their pharmaceutically acceptable salts, which can be derived from various organic and inorganic acids and bases by procedures known in the art. Pharmaceutically acceptable salt forms of the compounds of the formula I are for the most part prepared by conventional methods. If the compound of the formula I contains a carboxyl

group, one of its suitable salts can be formed by reacting the compound with a suitable base to give the corresponding base-addition salt. Such bases are, for example, alkali metal hydroxides, including potassium hydroxide, sodium hydroxide and lithium hydroxide; alkaline earth metal hydroxides, such as barium hydroxide and calcium hydroxide; alkali metal alkoxides, for example potassium ethoxide and sodium propoxide; and various organic bases, such as piperidine, diethanolamine and N-methyl-glutamine. The aluminium salts of the compounds of the formula I are likewise included. In the case of certain compounds of the formula I, acid-addition salts can be formed by treating these compounds with pharmaceutically acceptable organic and inorganic acids, for example hydrogen halides, such as hydrogen chloride, hydrogen bromide or hydrogen iodide, other mineral acids and corresponding salts thereof, such as sulfate, nitrate or phosphate and the like, and alkyl- and monoarylsulfonates, such as ethanesulfonate, toluenesulfonate and benzenesulfonate, and other organic acids and corresponding salts thereof, such as acetate, trifluoroacetate, tartrate, maleate, succinate, citrate, benzoate, salicylate, ascorbate and the like. Accordingly, pharmaceutically acceptable acid-addition salts of the compounds of the formula I include the following: acetate, adipate, alginate, arginate, aspartate, benzoate, benzenesulfonate (besylate), bisulfate, bisulfite, bromide, butyrate, camphorate, camphorsulfonate, caprylate, chloride, chlorobenzoate, citrate, cyclopentanepropionate, diglucconate, dihydrogenphosphate, dinitrobenzoate, dodecylsulfate, ethanesulfonate, fumarate, galacterate (from mucic acid), galacturonate, glucoheptanoate, gluconate, glutamate, glycerophosphate, hemisuccinate, hemisulfate, heptanoate, hexanoate, hippurate, hydrochloride, hydrobromide, hydroiodide, 2-hydroxyethanesulfonate, iodide, isethionate, isobutyrate, lactate, lactobionate, malate, maleate, malonate, mandelate, metaphosphate, methanesulfonate, methylbenzoate, monohydrogenphosphate, 2-naphthalenesulfonate, nicotinate, nitrate, oxalate, oleate, palmoate, pectinate, persulfate, phenylacetate, 3-phenylpropionate, phosphate, phosphonate, phthalate, but this does not represent a restriction.

Furthermore, the base salts of the compounds according to the invention include aluminium, ammonium, calcium, copper, iron(III), iron(II), lithium, magnesium, manganese(III), manganese(II), potassium, sodium and zinc salts, but this is not intended to represent a restriction. Of the above-mentioned salts, preference is given to ammonium; the alkali metal salts sodium and potassium, and the alkaline earth metal salts calcium and magnesium. Salts of the compounds of the formula I which are derived from pharmaceutically acceptable organic non-toxic bases include salts of primary, secondary and tertiary amines, substituted amines, also including naturally occurring substituted amines, cyclic amines, and basic ion exchanger resins, for example arginine, betaine, caffeine, chloroprocaine, choline, N,N'-dibenzylethylenediamine (benzathine), dicyclohexylamine, diethanolamine, diethylamine, 2-diethylaminoethanol, 2-dimethylaminoethanol, ethanolamine, ethylenediamine, N-ethylmorpholine, N-ethylpiperidine, glucamine, glucosamine, histidine, hydrabamine, isopropylamine, lidocaine, lysine, meglumine, N-methyl-D-glucamine, morpholine, piperazine, piperidine, polyamine resins, procaine, purines, theobromine, triethanolamine, triethylamine, trimethylamine, tripropylamine and tris(hydroxymethyl)methylamine (tromethamine), but this is not intended to represent a restriction.

Compounds of the present invention which contain basic nitrogen-containing groups can be quaternised using agents such as (C₁-C₄)alkyl halides, for example methyl, ethyl, isopropyl and tert-butyl chloride, bromide and iodide; di(C₁-C₄)alkyl sulfates, for example dimethyl, diethyl and diamyl sulfate; (C₁₀-C₁₈)alkyl halides, for example decyl, dodecyl, lauryl, myristyl and stearyl chloride, bromide and iodide; and aryl(C₁-C₄)alkyl halides, for example benzyl chloride and phenethyl bromide. Both water- and oil-soluble compounds according to the invention can be prepared using such salts.

The above-mentioned pharmaceutical salts which are preferred include acetate, trifluoroacetate, besylate, citrate, fumarate, gluconate, hemisuccinate, hippurate, hydrochloride, hydrobromide, isethionate, mandelate, meglumine, nitrate, oleate, phosphonate, pivalate, sodium phosphate, stearate, sulfate, sulfosalicylate, tartrate, thiomalate, tosylate and tromethamine, but this is not intended to represent a restriction.

The acid-addition salts of basic compounds of the formula I are prepared by bringing the free base form into contact with a sufficient amount of the desired acid, causing the formation of the salt in a conventional manner. The free base can be regenerated by bringing the salt form into contact with a base and isolating the free base in a conventional manner. The free base forms differ in a certain respect from the corresponding salt forms thereof with respect to certain physical properties, such as solubility in polar solvents; for the purposes of the invention, however, the salts otherwise correspond to the respective free base forms thereof.

As mentioned, the pharmaceutically acceptable base-addition salts of the compounds of the formula I are formed with metals or amines, such as alkali metals and alkaline earth metals or organic amines. Preferred metals are sodium, potassium, magnesium and calcium. Preferred organic amines are N,N'-dibenzylethylenediamine, chloroprocaine, choline, diethanolamine, ethylenediamine, N-methyl-D-glucamine and procaine.

The base-addition salts of acidic compounds according to the invention are prepared by bringing the free acid form into contact with a sufficient amount of the desired base, causing the formation of the salt in a conventional manner. The free acid can be regenerated by bringing the salt form into contact with an acid and isolating the free acid in a conventional manner. The free acid forms differ in a certain respect from the corresponding salt forms thereof with respect to certain physical properties, such as solubility

in polar solvents; for the purposes of the invention, however, the salts otherwise correspond to the respective free acid forms thereof.

If a compound according to the invention contains more than one group which is capable of forming pharmaceutically acceptable salts of this type, the invention also encompasses multiple salts. Typical multiple salt forms include, for example, bitartrate, diacetate, difumarate, dimeglumine, diphosphate, disodium and trihydrochloride, but this is not intended to represent a restriction.

With regard to that stated above, it can be seen that the term "pharmaceutically acceptable salt" in the present connection is taken to mean an active ingredient which comprises a compound of the formula I in the form of one of its salts, in particular if this salt form imparts improved pharmacokinetic properties on the active ingredient compared with the free form of the active ingredient or any other salt form of the active ingredient used earlier. The pharmaceutically acceptable salt form of the active ingredient can also provide this active ingredient for the first time with a desired pharmacokinetic property which it did not have earlier and can even have a positive influence on the pharmacodynamics of this active ingredient with respect to its therapeutic efficacy in the body.

The invention furthermore relates to medicaments comprising at least one compound of the formula I and/or pharmaceutically usable derivatives, solvates and stereoisomers thereof, including mixtures thereof in all ratios, and optionally excipients and/or adjuvants.

Pharmaceutical formulations can be administered in the form of dosage units which comprise a predetermined amount of active ingredient per dosage unit. Such a unit can comprise, for example, 0.5 mg to 1 g, preferably 1 mg to 700 mg, particularly preferably 5 mg to 100 mg, of a compound according to the invention, depending on the condition treated, the method of

administration and the age, weight and condition of the patient, or pharmaceutical formulations can be administered in the form of dosage units which comprise a predetermined amount of active ingredient per dosage unit. Preferred dosage unit formulations are those which comprise a daily dose or part-dose, as indicated above, or a corresponding fraction thereof of an active ingredient. Furthermore, pharmaceutical formulations of this type can be prepared using a process which is generally known in the pharmaceutical art.

Pharmaceutical formulations can be adapted for administration via any desired suitable method, for example by oral (including buccal or sublingual), rectal, nasal, topical (including buccal, sublingual or transdermal), vaginal or parenteral (including subcutaneous, intramuscular, intravenous or intra-dermal) methods. Such formulations can be prepared using all processes known in the pharmaceutical art by, for example, combining the active ingredient with the excipient(s) or adjuvant(s).

Pharmaceutical formulations adapted for oral administration can be administered as separate units, such as, for example, capsules or tablets; powders or granules; solutions or suspensions in aqueous or non-aqueous liquids; edible foams or foam foods; or oil-in-water liquid emulsions or water-in-oil liquid emulsions.

Thus, for example, in the case of oral administration in the form of a tablet or capsule, the active-ingredient component can be combined with an oral, non-toxic and pharmaceutically acceptable inert excipient, such as, for example, ethanol, glycerol, water and the like. Powders are prepared by comminuting the compound to a suitable fine size and mixing it with a pharmaceutical excipient comminuted in a similar manner, such as, for example, an edible carbohydrate, such as, for example, starch or mannitol. A flavour, preservative, dispersant and dye may likewise be present.

Capsules are produced by preparing a powder mixture as described above and filling shaped gelatine shells therewith. Glidants and lubricants, such as, for example, highly disperse silicic acid, talc, magnesium stearate, calcium stearate or polyethylene glycol in solid form, can be added to the powder mixture before the filling operation. A disintegrant or solubiliser, such as, for example, agar-agar, calcium carbonate or sodium carbonate, may likewise be added in order to improve the availability of the medicament after the capsule has been taken.

In addition, if desired or necessary, suitable binders, lubricants and disintegrants as well as dyes can likewise be incorporated into the mixture. Suitable binders include starch, gelatine, natural sugars, such as, for example, glucose or beta-lactose, sweeteners made from maize, natural and synthetic rubber, such as, for example, acacia, tragacanth or sodium alginate, carboxymethylcellulose, polyethylene glycol, waxes, and the like. The lubricants used in these dosage forms include sodium oleate, sodium stearate, magnesium stearate, sodium benzoate, sodium acetate, sodium chloride and the like. The disintegrants include, without being restricted thereto, starch, methylcellulose, agar, bentonite, xanthan gum and the like. The tablets are formulated by, for example, preparing a powder mixture, granulating or dry-pressing the mixture, adding a lubricant and a disintegrant and pressing the entire mixture to give tablets. A powder mixture is prepared by mixing the compound comminuted in a suitable manner with a diluent or a base, as described above, and optionally with a binder, such as, for example, carboxymethylcellulose, an alginate, gelatine or polyvinylpyrrolidone, a dissolution retardant, such as, for example, paraffin, an absorption accelerator, such as, for example, a quaternary salt, and/or an absorbant, such as, for example, bentonite, kaolin or dicalcium phosphate. The powder mixture can be granulated by wetting it with a binder, such as, for example, syrup, starch paste, acadia mucilage or solutions of cellulose or polymer materials and pressing it through a sieve. As an alternative to

granulation, the powder mixture can be run through a tabletting machine, giving lumps of non-uniform shape which are broken up to form granules. The granules can be lubricated by addition of stearic acid, a stearate salt, talc or mineral oil in order to prevent sticking to the tablet casting moulds. The lubricated mixture is then pressed to give tablets. The compounds according to the invention can also be combined with a free-flowing inert excipient and then pressed directly to give tablets without carrying out the granulation or dry-pressing steps. A transparent or opaque protective layer consisting of a shellac sealing layer, a layer of sugar or polymer material and a gloss layer of wax may be present. Dyes can be added to these coatings in order to be able to differentiate between different dosage units.

Oral liquids, such as, for example, solution, syrups and elixirs, can be prepared in the form of dosage units so that a given quantity comprises a pre-specified amount of the compound. Syrups can be prepared by dissolving the compound in an aqueous solution with a suitable flavour, while elixirs are prepared using a non-toxic alcoholic vehicle. Suspensions can be formulated by dispersion of the compound in a non-toxic vehicle. Solubilisers and emulsifiers, such as, for example, ethoxylated isostearyl alcohols and polyoxyethylene sorbitol ethers, preservatives, flavour additives, such as, for example, peppermint oil or natural sweeteners or saccharin, or other artificial sweeteners and the like, can likewise be added.

The dosage unit formulations for oral administration can, if desired, be encapsulated in microcapsules. The formulation can also be prepared in such a way that the release is extended or retarded, such as, for example, by coating or embedding of particulate material in polymers, wax and the like.

The compounds of the formula I and salts, solvates and physiologically functional derivatives thereof can also be administered in the form of liposome delivery systems, such as, for example, small unilamellar vesicles,

large unilamellar vesicles and multilamellar vesicles. Liposomes can be formed from various phospholipids, such as, for example, cholesterol, stearylamine or phosphatidylcholines.

The compounds of the formula I and the salts, solvates and physiologically functional derivatives thereof can also be delivered using monoclonal antibodies as individual carriers to which the compound molecules are coupled. The compounds can also be coupled to soluble polymers as targeted medicament carriers. Such polymers may encompass polyvinylpyrrolidone, pyran copolymer, polyhydroxypropylmethacrylamidophenol, polyhydroxyethylaspartamidophenol or polyethylene oxide polylysine, substituted by palmitoyl radicals. The compounds may furthermore be coupled to a class of biodegradable polymers which are suitable for achieving controlled release of a medicament, for example polylactic acid, poly-epsilon-caprolactone, polyhydroxybutyric acid, polyorthoesters, polyacetals, polydihydroxypyrans, polycyanoacrylates and crosslinked or amphipathic block copolymers of hydrogels.

Pharmaceutical formulations adapted for transdermal administration can be administered as independent plasters for extended, close contact with the epidermis of the recipient. Thus, for example, the active ingredient can be delivered from the plaster by iontophoresis, as described in general terms in *Pharmaceutical Research*, 3(6), 318 (1986).

Pharmaceutical compounds adapted for topical administration can be formulated as ointments, creams, suspensions, lotions, powders, solutions, pastes, gels, sprays, aerosols or oils.

For the treatment of the eye or other external tissue, for example mouth and skin, the formulations are preferably applied as topical ointment or cream. In the case of formulation to give an ointment, the active ingredient can be employed either with a paraffinic or a water-miscible cream base.

Alternatively, the active ingredient can be formulated to give a cream with an oil-in-water cream base or a water-in-oil base.

Pharmaceutical formulations adapted for topical application to the eye include eye drops, in which the active ingredient is dissolved or suspended in a suitable carrier, in particular an aqueous solvent.

Pharmaceutical formulations adapted for topical application in the mouth encompass lozenges, pastilles and mouthwashes.

Pharmaceutical formulations adapted for rectal administration can be administered in the form of suppositories or enemas.

Pharmaceutical formulations adapted for nasal administration in which the carrier substance is a solid comprise a coarse powder having a particle size, for example, in the range 20-500 microns, which is administered in the manner in which snuff is taken, i.e. by rapid inhalation via the nasal passages from a container containing the powder held close to the nose. Suitable formulations for administration as nasal spray or nose drops with a liquid as carrier substance encompass active-ingredient solutions in water or oil.

Pharmaceutical formulations adapted for administration by inhalation encompass finely particulate dusts or mists, which can be generated by various types of pressurised dispensers with aerosols, nebulisers or insufflators.

Pharmaceutical formulations adapted for vaginal administration can be administered as pessaries, tampons, creams, gels, pastes, foams or spray formulations.

Pharmaceutical formulations adapted for parenteral administration include aqueous and non-aqueous sterile injection solutions comprising antioxidants, buffers, bacteriostatics and solutes, by means of which the formulation is rendered isotonic with the blood of the recipient to be treated; and aqueous and non-aqueous sterile suspensions, which may comprise suspension media and thickeners. The formulations can be administered in single-dose or multidose containers, for example sealed ampoules and vials, and stored in freeze-dried (lyophilised) state, so that only the addition of the sterile carrier liquid, for example water for injection purposes, immediately before use is necessary.

Injection solutions and suspensions prepared in accordance with the recipe can be prepared from sterile powders, granules and tablets.

It goes without saying that, in addition to the above particularly mentioned constituents, the formulations may also comprise other agents usual in the art with respect to the particular type of formulation; thus, for example, formulations which are suitable for oral administration may comprise flavours.

A therapeutically effective amount of a compound of the formula I depends on a number of factors, including, for example, the age and weight of the animal, the precise condition which requires treatment, and its severity, the nature of the formulation and the method of administration, and is ultimately determined by the treating doctor or vet. However, an effective amount of a compound according to the invention for the treatment of neoplastic growth, for example colon or breast carcinoma, is generally in the range from 0.1 to 100 mg/kg of body weight of the recipient (mammal) per day and particularly typically in the range from 1 to 10 mg/kg of body weight per day. Thus, the actual amount per day for an adult mammal weighing 70 kg is usually between 70 and 700 mg, where this amount can be administered as a single dose per day or usually in a series of part-doses (such as, for example, two, three, four, five or six) per day, so that the total daily dose is the same.

An effective amount of a salt or solvate or of a physiologically functional derivative thereof can be determined as the fraction of the effective amount of the compound according to the invention *per se*. It can be assumed that similar doses are suitable for the treatment of other conditions mentioned above.

The invention furthermore relates to medicaments comprising at least one compound of the formula I and/or pharmaceutically usable derivatives, solvates and stereoisomers thereof, including mixtures thereof in all ratios, and at least one further medicament active ingredient.

The invention also relates to a set (kit) consisting of separate packs of

- (a) an effective amount of a compound of the formula I and/or pharmaceutically usable derivatives, solvates and stereoisomers thereof, including mixtures thereof in all ratios,
and
- (b) an effective amount of a further medicament active ingredient.

The set comprises suitable containers, such as boxes, individual bottles, bags or ampoules. The set may, for example, comprise separate ampoules, each containing an effective amount of a compound of the formula I and/or pharmaceutically usable derivatives, solvates and stereoisomers thereof, including mixtures thereof in all ratios,
and an effective amount of a further medicament active ingredient in dissolved or lyophilised form.

The medicaments from Table 1 are preferably, but not exclusively, combined with the compounds of the formula I. A combination of the formula I and medicaments from Table 1 can also be combined with compounds of the formula V.

Table 1.

Alkylating agents	Cyclophosphamide	Lomustine
	Busulfan	Procarbazine
	Ifosfamide	Altretamine
	Melphalan	Estramustine phosphate
	Hexamethylmelamine	Mechloroethamine
	Thiotepa	Streptozocin
	Chloroambucil	Temozolamide
	Dacarbazine	Semustine
	Carmustine	
Platinum agents	Cisplatin	Carboplatin
	Oxaliplatin	ZD-0473 (AnorMED)
	Spiroplatin	Lobaplatin (Aetema)
	Carboxyphthalatoplatinum	Satraplatin (Johnson Matthey)
	Tetraplatin	BBR-3464
	Orniplatin	(Hoffmann-La Roche)
	Iproplatin	SM-11355 (Sumitomo)
		AP-5280 (Access)
Antimetabolites	Azacytidine	Tomudex
	Gemcitabine	Trimetrexate
	Capecitabine	Deoxycytidine
	5-Fluorouracil	Fludarabine
	Floxuridine	Pentostatin
	2-Chlorodesoxyadenosine	Raltitrexed
	6-Mercaptopurine	Hydroxyurea
	6-Thioguanine	Decitabine (SuperGen)
	Cytarabine	Clofarabine (Bioenvision)
	2-Fluorodesoxycytidine	Irofulven (MGI Pharma)
	Methotrexate	DMDC (Hoffmann-La Roche)
	Idarubicin	Ethylnucleotide (Taiho)
Topoisomerase inhibitors	Amsacrine	Rubitecan (SuperGen)
	Epirubicin	Exatecan mesylate
	Etoposide	(Daiichi)
	Teniposide or mitoxantrone	Quinamed (ChemGenex)
	Irinotecan (CPT-11)	Gimatecan (Sigma-Tau)
	7-Ethyl-10-hydroxycamptothecin	Diflomotecan (Beaufour-Ipsen)
	Topotecan	TAS-103 (Taiho)
	Dexrazoxane (TopoTarget)	Elsamitrucin (Spectrum)
	Pixantrone (Novuspharma)	J-107088 (Merck & Co)
		BNP-1350 (BioNumerik)
		CKD-602 (Chong Kun)

	Rebeccamycin analogue (Exelixis) BBR-3576 (Novuspharrna)	Dang) KW-2170 (Kyowa Hakko)
Antitumour antibiotics	Dactinomycin (Actinomycin D) Doxorubicin (Adriamycin) Deoxyrubicin Valrubicin Daunorubicin (Daunomycin) Epirubicin Therarubicin Idarubicin Rubidazon Plicamycinp Porfiromycin Cyanomorpholinodoxorubicin Mitoxantron (Novantron)	Amonafide Azonaflde Anthrapyrazole Oxantrazole Losoxantrone Bleomycin sulfate (Blenoxan) Bleomycinic acid Bleomycin A Bleomycin B Mitomycin C MEN-10755 (Menarini) GPX-100 (Gem Pharmaceuticals)
Antimitotic agents	Paclitaxel Docetaxel Colchicine Vinblastine Vincristine Vinorelbine Vindesine Dolastatin 10 (NCI) Rhizoxin (Fujisawa) Mivobulin (Warner-Lambert) Cemadotin (BASF) RPR 109881A (Aventis) TXD 258 (Aventis) Epothilone B (Novartis) T 900607 (Tularik) T 138067 (Tularik) Cryptophycin 52 (Eli Lilly) Vinflunine (Fabre) Auristatin PE (Teikoku Hormone) BMS 247550 (BMS) BMS 184476 (BMS) BMS 188797 (BMS) Taxoprexin (Protarga)	SB 408075 (GlaxoSmithKline) E7010 (Abbott) PG-TXL (Cell Therapeutics) IDN 5109 (Bayer) A 105972 (Abbott) A 204197 (Abbott) LU 223651 (BASF) D 24851 (ASTA Medica) ER-86526 (Eisai) Combretastatin A4 (BMS) Isohomohalichondrin-B (PharmaMar) ZD 6126 (AstraZeneca) PEG-Paclitaxel (Enzon) AZ10992 (Asahi) IDN-5109 (Indena) AVLB (Prescient NeuroPharma) Azaepothilon B (BMS) BNP- 7787 (BioNumerik) CA-4-Prodrug (OXiGENE) Dolastatin-10 (NrH) CA-4 (OXiGENE)

Aromatase inhibitors	Aminoglutethimide Letrozole Anastrazole Formestan	Exemestan Atamestan (BioMedicines) YM-511 (Yamanouchi)
Thymidylate synthase inhibitors	Pemetrexed (Eli Lilly) ZD-9331 (BTG)	Nolatrexed (Eximias) CoFactor™ (BioKeys)
DNA antagonists	Trabectedin (PharmaMar) Glufosfamide (Baxter International) Albumin + 32P (Isotope Solutions) Thymectacin (NewBiotics) Edotreotid (Novartis)	Mafosfamide (Baxter International) Apaziquone (Spectrum Pharmaceuticals) O6-Benzylguanine (Paligent)
Farnesyl transferase inhibitors	Arglabin (NuOncology Labs) Ionafarnib (Schering-Plough) BAY-43-9006 (Bayer)	Tipifarnib (Johnson & Johnson) Perillyl alcohol (DOR BioPharma)
Pump inhibitors	CBT-1 (CBA Pharma) Tariquidar (Xenova) MS-209 (Schering AG)	Zosuquidar trihydrochloride (Eli Lilly) Biricodar dicitrate (Vertex)
Histone acetyl transferase inhibitors	Tacedinaline (Pfizer) SAHA (Aton Pharma) MS-275 (Schering AG)	Pivaloyloxymethyl butyrate (Titan) Depsi peptide (Fujisawa)
Metalloproteinase inhibitors Ribonucleoside reductase inhibitors	Neovastat (Aeterna Laboratories) Marimastat (British Biotech) Gallium maltolato (Titan) Triapin (Vion)	CMT -3 (CollaGenex) BMS-275291 (Celltech) Tezacitabine (Aventis) Didox (Molecules for Health)
TNF-alpha agonists/ antagonists	Virulizin (Lorus Therapeutics) CDC-394 (Celgene)	Revimid (Celgene)
Endothelin-A receptor antagonists	Atrasentan (Abbot) ZD-4054 (AstraZeneca)	YM-598 (Yamanouchi)

Retinoic acid receptor agonists	Fenretinide (Johnson & Johnson) LGD-1550 (Ligand)	Alitretinoin (Ligand)
Immuno-modulators	Interferon Oncophage (Antigenics) GMK (Progenics) Adenocarcinoma vaccine (Biomira) CTP-37 (AVI BioPharma) JRX-2 (Immuno-Rx) PEP-005 (Peplin Biotech) Synchrovax vaccines (CTL Immuno) Melanoma vaccine (CTL Immuno) p21-RAS vaccine (GemVax)	Dexosome therapy (Anosys) Pentrix (Australian Cancer Technology) JSF-154 (Tragen) Cancer vaccine (Intercell) Norelin (Biostar) BLP-25 (Biomira) MGV (Progenics) !3-Alethin (Dovetail) CLL-Thera (Vasogen)
Hormonal and antihormonal agents	Oestrogens Conjugated oestrogens Ethynodiol Chlorotrianisene Idenestrol Hydroxyprogesterone caproate Medroxyprogesterone Testosterone Testosterone propionate Fluoxymesterone Methyltestosterone Diethylstilbestrol Megestrol Tamoxifen Toremofin Dexamethasone	Prednisone Methylprednisolone Prednisolone Aminoglutethimide Leuprorelin Goserelin Leuporelin Bicalutamide Flutamide Octreotide Nilutamide Mitotan P-04 (Novogen) 2-Methoxyoestradiol (EntreMed) Arzoxifен (Eli Lilly)
Photodynamic agents	Talaporfin (Light Sciences) Theralux (Theratechnologies) Motexafin-Gadolinium (Pharmacyclics)	Pd-Bacteriopheophorbid (Yeda) Lutetium-Texaphyrin (Pharmacyclics) Hypericin
Tyrosine kinase inhibitors	Imatinib (Novartis) Leflunomide(Sugen/Pharmacia) ZDI839 (AstraZeneca)	Kahalide F (PharmaMar) CEP- 701 (Cephalon) CEP-751 (Cephalon) MLN518 (Millenium)

	Erlotinib (Oncogene Science) Canertjnib (Pfizer) Squalamine (Genaera) SU5416 (Pharmacia) SU6668 (Pharmacia) ZD4190 (AstraZeneca) ZD6474 (AstraZeneca) Vatalanib (Novartis) PKI166 (Novartis) GW2016 (GlaxoSmithKline) EKB-509 (Wyeth) EKB-569 (Wyeth)	PKC412 (Novartis) Phenoxodiol O Trastuzumab (Genentech) C225 (ImClone) rhu-Mab (Genentech) MDX-H210 (Medarex) 2C4 (Genentech) MDX-447 (Medarex) ABX-EGF (Abgenix) IMC-1C11 (ImClone)
Various agents	SR-27897 (CCK-A inhibitor, Sanofi-Synthelabo) Tocladesine (cyclic AMP agonist, Ribapharm) Alvocidib (CDK inhibitor, Aventis) CV-247 (COX-2 inhibitor, Ivy Medical) P54 (COX-2 inhibitor, Phytopharm) CapCell™ (CYP450 stimulant, Bavarian Nordic) GCS-IOO (gal3 antagonist, GlycoGenesys) G17DT immunogen (gastrin inhibitor, Aphanton) Efaproxiral (oxygenator, Allos Therapeutics) PI-88 (heparanase inhibitor, Progen) Tesmilifene (histamine antagonist, YM BioSciences) Histamine (histamine H2 receptor agonist, Maxim) Tiazofurin (IMPDH inhibitor, Ribapharm) Cilengitide (integrin antagonist, Merck KGaA) SR-31747 (IL-1 antagonist, Sanofi-Synthelabo) CCI-779 (mTOR kinase	BCX-1777 (PNP inhibitor, BioCryst) Ranpirnase (ribonuclease stimulant, Alfacell) Galarubicin (RNA synthesis inhibitor, Dong-A) Tirapazamine (reducing agent, SRI International) N-Acetylcysteine (reducing agent, Zambon) R-Flurbiprofen (NF-kappaB inhibitor, Encore) 3CPA (NF-kappaB inhibitor, Active Biotech) Seocalcitrol (vitamin D receptor agonist, Leo) 131-I-TM-601 (DNA antagonist, TransMolecular) Eflornithin (ODC inhibitor, ILEX Oncology) Minodronic acid (osteoclast inhibitor, Yamanouchi) Indisulam (p53 stimulant, Eisai) Aplidin (PPT inhibitor, PharmaMar) Rituximab (CD20 antibody, Genentech) Gemtuzumab (CD33 antibody, Wyeth Ayerst)

	inhibitor, Wyeth) Exisulind (PDE-V inhibitor, Cell Pathways) CP-461 (PDE-V inhibitor, Cell Pathways) AG-2037 (GART inhibitor, Pfizer) WX-UK1 (plasminogen activator inhibitor, Wilex) PBI-1402 (PMN stimulant, ProMetic LifeSciences) Bortezomib (proteasome inhibitor, Millennium) SRL-172 (T-cell stimulant, SR Pharma) TLK-286 (glutathione-S transferase inhibitor, Telik) PT-100 (growth factor agonist, Point Therapeutics) Midostaurin (PKC inhibitor, Novartis) Bryostatin-1 (PKC stimulant, GPC Biotech) CDA-II (apoptosis promoter, Everlife) SDX-101 (apoptosis promoter, Salmedix) Ceftafotin (apoptosis promoter, ChemGenex)	PG2 (haematopoiesis promoter, Pharmagenesis) Immunol™ (triclosan mouthwash, Endo) Triacetyluridine (uridine prodrug, Wellstat) SN-4071 (sarcoma agent, Signature BioScience) TransMID-107™ (immunotoxin, KS Biomedix) PCK-3145 (apoptosis promoter, Procyon) Doranidazole (apoptosis promoter, Pola) CHS-828 (cytotoxic agent, Leo) Trans-retinic acid (differentiator, NIH) MX6 (apoptosis promoter, MAXIA) Apomine (apoptosis promoter, ILEX Oncology) Urocidin (apoptosis promoter, Bioniche) Ro-31-7453 (apoptosis promoter, La Roche) Brostallicin (apoptosis promoter, Pharmacia)
Alkylating agents	Cyclophosphamide Busulfan Ifosfamide Melphalan Hexamethylmelamine Thiotepa Chloroambucil Dacarbazine Carmustine	Lomustine Procarbazine Altretamine Estramustine phosphate Mechloroethamine Streptozocin Temozolamide Semustine
Platinum agents	Cisplatin Oxaliplatin Spiroplatin Carboxyphthalatoplatinum Tetraplatin Ormiplatin	Carboplatin ZD-0473 (AnorMED) Lobaplatin (Aetema) Satraplatin (Johnson Matthey) BBR-3464

	Iproplatin	(Hoffmann-La Roche) SM-11355 (Sumitomo) AP-5280 (Access)
Antimetabolites	Azacytidine Gemcitabine Capecitabine 5-Fluorouracil Floxuridine 2-Chlorodesoxyadenosine 6-Mercaptopurine 6-Thioguanine Cytarabine 2-Fluorodesoxycytidine Methotrexate Idatrexate	Tomudex Trimetrexate Deoxycoformycin Fludarabine Pentostatin Raltitrexed Hydroxyurea Decitabine (SuperGen) Clofarabine (Bioenvision) Irofulven (MGI Pharma) DMDC (Hoffmann-La Roche) Ethynylcytidine (Taiho)
Topoisomerase inhibitors	Amsacrine Epirubicin Etoposide Teniposide or mitoxantrone Irinotecan (CPT-11) 7-Ethyl-10-hydroxycamptothecin Topotecan Dexrazoxane (TopoTarget) Pixantrone (Novuspharma) Rebeccamycin analogue (Exelixis) BBR-3576 (Novuspharma)	Rubitecan (SuperGen) Exatecan mesylate (Daiichi) Quinamed (ChemGenex) Gimatecan (Sigma-Tau) Diflomotecan (Beaufour-Ipsen) TAS-103 (Taiho) Elsamitrucin (Spectrum) J-107088 (Merck & Co) BNP-1350 (BioNumerik) CKD-602 (Chong Kun Dang) KW-2170 (Kyowa Hakko)
Antitumour antibiotics	Dactinomycin (Actinomycin D) Doxorubicin (Adriamycin) Deoxyrubicin Valrubicin Daunorubicin (Daunomycin) Epirubicin Therarubicin Idarubicin Rubidazon Plicamycin Porfiromycin	Amonafide Azonafide Anthrapyrazole Oxantrazole Losoxantrone Bleomycin sulfate (Blenoxan) Bleomycinic acid Bleomycin A Bleomycin B Mitomycin C MEN-10755 (Menarini) GPX-100 (Gem)

	Cyanomorpholinodoxo-rubicin Mitoxantron (Novantron)	Pharmaceuticals)
Antimitotic agents	Paclitaxel Docetaxel Colchicine Vinblastine Vincristine Vinorelbine Vindesine Dolastatin 10 (NCI) Rhizoxin (Fujisawa) Mivobulin (Warner-Lambert) Cemadotin (BASF) RPR 109881A (Aventis) TXD 258 (Aventis) Epothilone B (Novartis) T 900607 (Tularik) T 138067 (Tularik) Cryptophycin 52 (Eli Lilly) Vinflunine (Fabre) Auristatin PE (Teikoku Hormone) BMS 247550 (BMS) BMS 184476 (BMS) BMS 188797 (BMS) Taxoprexin (Protarga)	SB 408075 (GlaxoSmithKline) E7010 (Abbott) PG-TXL (Cell Therapeutics) IDN 5109 (Bayer) A 105972 (Abbott) A 204197 (Abbott) LU 223651 (BASF) D 24851 (ASTA Medica) ER-86526 (Eisai) Combretastatin A4 (BMS) Isohomohalichondrin-B (PharmaMar) ZD 6126 (AstraZeneca) PEG-Paclitaxel (Enzon) AZ10992 (Asahi) !DN-5109 (Indena) AVLB (Prescient NeuroPharma) Azaepothilon B (BMS) BNP- 7787 (BioNumerik) CA-4-Prodrug (OXiGENE) Dolastatin-10 (NrH) CA-4 (OXiGENE)
Aromatase inhibitors	Aminoglutethimide Letrozole Anastrazole Formestan	Exemestan Atamestan (BioMedicines) YM-511 (Yamanouchi)
Thymidylate synthase inhibitors	Pemetrexed (Eli Lilly) ZD-9331 (BTG)	Nolatrexed (Eximias) CoFactor™ (BioKeys)
DNA antagonists	Trabectedin (PharmaMar) Glufosfamide (Baxter International) Albumin + 32P (Isotope Solutions) Thymectacin (NewBiotics) Edotreotide (Novartis)	Mafosfamide (Baxter International) Apaziquone (Spectrum Pharmaceuticals) O6-Benzylguanine (Palgent)

Farnesyl transferase inhibitors	Arglabin (NuOncology Labs) Ionaflarnib (Schering-Plough) BAY-43-9006 (Bayer)	Tipifarnib (Johnson & Johnson) Perillyl alcohol (DOR BioPharma)
Pump inhibitors	CBT-1 (CBA Pharma) Tariquidar (Xenova) MS-209 (Schering AG)	Zosuquidar trihydrochloride (Eli Lilly) Biricodar dicitrate (Vertex)
Histone acetyl transferase inhibitors	Tacedinaline (Pfizer) SAHA (Aton Pharma) MS-275 (Schering AG)	Pivaloyloxymethyl butyrate (Titan) Depsi peptide (Fujisawa)
Metalloproteinase inhibitors Ribonucleoside reductase inhibitors	Neovastat (Aeterna Laboratories) Marimastat (British Biotech) Gallium maltolate (Titan) Triapin (Vion)	CMT -3 (CollaGenex) BMS-275291 (Celltech) Tezacitabine (Aventis) Didox (Molecules for Health)
TNF-alpha agonists/ antagonists	Virulizin (Lorus Therapeutics) CDC-394 (Celgene)	Revimid (Celgene)
Endothelin-A receptor antagonists	Atrasentan (Abbot) ZD-4054 (AstraZeneca)	YM-598 (Yamanouchi)
Retinoic acid receptor agonists	Fenretinide (Johnson & Johnson) LGD-1550 (Ligand)	Alitretinoin (Ligand)
Immuno-modulators	Interferon Oncophage (Antigenics) GMK (Progenics) Adenocarcinoma vaccine (Biomira) CTP-37 (AVI BioPharma) JRX-2 (Immuno-Rx) PEP-005 (Peplin Biotech) Synchrovax vaccines (CTL Immuno) Melanoma vaccine (CTL Immuno) p21-RAS vaccine (GemVax)	Dexosome therapy (Anosys) Pentrix (Australian Cancer Technology) JSF-154 (Tragen) Cancer vaccine (Intercell) Norelin (Biostar) BLP-25 (Biomira) MGV (Progenics) I3-Alethin (Dovetail) CLL-Thera (Vasogen)

Hormonal and antihormonal agents	Oestrogens Conjugated oestrogens Ethynyloestradiol Chlorotrianisene Idenestrol Hydroxyprogesterone caproate Medroxyprogesterone Testosterone Testosterone propionate Fluoxymesterone Methyltestosterone Diethylstilbestrol Megestrol Tamoxifen Toremofin Dexamethasone	Prednisone Methylprednisolone Prednisolone Aminoglutethimide Leuprolide Goserelin Leuporelin Bicalutamide Flutamide Octreotide Nilutamide Mitotan P-04 (Novogen) 2-Methoxyoestradiol (EntreMed) Arzoxifen (Eli Lilly)
Photodynamic agents	Talaporfin (Light Sciences) Theralux (Theratechnologies) Motexafin-Gadolinium (Pharmacyclicks)	Pd-Bacteriopheophorbid (Yeda) Lutetium-Texaphyrin (Pharmacyclicks) Hypericin
Tyrosine kinase inhibitors	Imatinib (Novartis) Leflunomide(Sugen/Pharmacia) ZDI839 (AstraZeneca) Erlotinib (Oncogene Science) Canertjnib (Pfizer) Squalamine (Genaera) SU5416 (Pharmacia) SU6668 (Pharmacia) ZD4190 (AstraZeneca) ZD6474 (AstraZeneca) Vatalanib (Novartis) PKI166 (Novartis) GW2016 (GlaxoSmithKline) EKB-509 (Wyeth) EKB-569 (Wyeth)	Kahalide F (PharmaMar) CEP- 701 (Cephalon) CEP-751 (Cephalon) MLN518 (Millenium) PKC412 (Novartis) Phenoxydiol O Trastuzumab (Genentech) C225 (ImClone) rhu-Mab (Genentech) MDX-H210 (Medarex) 2C4 (Genentech) MDX-447 (Medarex) ABX-EGF (Abgenix) IMC-1C11 (ImClone)
Various agents	SR-27897 (CCK-A inhibitor, Sanofi-Synthelabo) Tocladesine (cyclic AMP	BCX-1777 (PNP inhibitor, BioCryst) Ranpirnase (ribonuclease stimulant, Alfacell)

	agonist, Ribapharm) Alvocidib (CDK inhibitor, Aventis) CV-247 (COX-2 inhibitor, Ivy Medical) P54 (COX-2 inhibitor, Phytopharm) CapCell™ (CYP450 stimulant, Bavarian Nordic) GCS-IOO (gal3 antagonist, GlycoGenesys) G17DT immunogen (gastrin inhibitor, Aphton) Efaproxiral (oxygenator, Allos Therapeutics) PI-88 (heparanase inhibitor, Progen) Tesmilifene (histamine antagonist, YM BioSciences) Histamine (histamine H2 receptor agonist, Maxim) Tiazofurin (IMPDH inhibitor, Ribapharm) Cilengitide (integrin antagonist, Merck KGaA) SR-31747 (IL-1 antagonist, Sanofi-Synthelabo) CCI-779 (mTOR kinase inhibitor, Wyeth) Exisulind (PDE-V inhibitor, Cell Pathways) CP-461 (PDE-V inhibitor, Cell Pathways) AG-2037 (GART inhibitor, Pfizer) WX-UK1 (plasminogen activator inhibitor, Wilex) PBI-1402 (PMN stimulant, ProMetic LifeSciences) Bortezomib (proteasome inhibitor, Millennium) SRL-172 (T-cell stimulant, SR Pharma) TLK-286 (glutathione-S transferase inhibitor, Telik) PT-100 (growth factor)	Galarubicin (RNA synthesis inhibitor, Dong-A) Tirapazamine (reducing agent, SRI International) N-Acetylcysteine (reducing agent, Zambon) R-Flurbiprofen (NF-kappaB inhibitor, Encore) 3CPA (NF-kappaB inhibitor, Active Biotech) Seocalcitol (vitamin D receptor agonist, Leo) 131-I-TM-601 (DNA antagonist, TransMolecular) Eflornithin (ODC inhibitor, ILEX Oncology) Minodronic acid (osteoclast inhibitor, Yamanouchi) Indisulam (p53 stimulant, Eisai) Aplidin (PPT inhibitor, PharmaMar) Rituximab (CD20 antibody, Genentech) Gemtuzumab (CD33 antibody, Wyeth Ayerst) PG2 (haematopoiesis promoter, Pharmagenesis) Immunol™ (triclosan mouthwash, Endo) Triacetyluridine (uridine prodrug, Wellstat) SN-4071 (sarcoma agent, Signature BioScience) TransMID-107™ (immunotoxin, KS Biomedix) PCK-3145 (apoptosis promoter, Procyon) Doranidazole (apoptosis promoter, Pola) CHS-828 (cytotoxic agent, Leo) Trans-retinic acid
--	---	--

	agonist, Point Therapeutics) Midostaurin (PKC inhibitor, Novartis) Bryostatin-1 (PKC stimulant, GPC Biotech) CDA-II (apoptosis promoter, Everlife) SDX-101 (apoptosis promoter, Salmedix) Ceflafonin (apoptosis promoter, ChemGenex)	(differentiator, NIH) MX6 (apoptosis promoter, MAXIA) Apomine (apoptosis promoter, ILEX Oncology) Urocidin (apoptosis promoter, Bioniche) Ro-31-7453 (apoptosis promoter, La Roche) Brostallicin (apoptosis promoter, Pharmacia)
--	---	---

The compounds of the formula I are preferably combined with known anti-cancer agents:

The present compounds are also suitable for combination with known anti-cancer agents. These known anti-cancer agents include the following: oestrogen receptor modulators, androgen receptor modulators, retinoid receptor modulators, cytotoxic agents, antiproliferative agents, prenyl-protein transferase inhibitors, HMG-CoA reductase inhibitors, HIV protease inhibitors, reverse transcriptase inhibitors and other angiogenesis inhibitors. The present compounds are particularly suitable for administration at the same time as radiotherapy. The synergistic effects of inhibition of VEGF in combination with radiotherapy have been described by specialists (see WO 00/61186).

"Oestrogen receptor modulators" refers to compounds which interfere with or inhibit the binding of oestrogen to the receptor, regardless of mechanism. Examples of oestrogen receptor modulators include, but are not limited to, tamoxifen, raloxifene, idoxifene, LY353381, LY 117081, toremifene, fulvestrant, 4-[7-(2,2-dimethyl-1-oxopropoxy-4-methyl-2-[4-[2-(1-piperidinyl)ethoxy]phenyl]-2H-1-benzopyran-3-yl]phenyl 2,2-dimethylpropanoate, 4,4'-dihydroxybenzophenone-2,4-dinitrophenylhydrazone and SH646.

"Androgen receptor modulators" refers to compounds which interfere with or inhibit the binding of androgens to the receptor, regardless of mechanism. Examples of androgen receptor modulators include finasteride and

other 5 α -reductase inhibitors, nilutamide, flutamide, bicalutamide, liarozole and abiraterone acetate.

"Retinoid receptor modulators" refers to compounds which interfere with or inhibit the binding of retinoids to the receptor, regardless of mechanism.

Examples of such retinoid receptor modulators include bexarotene, tretinoin, 13-cis-retinoic acid, 9-cis-retinoic acid, α -difluoromethylornithine, ILX23-7553, trans-N-(4'-hydroxyphenyl)retinamide and N-4-carboxyphenyl-retinamide.

"Cytotoxic agents" refers to compounds which result in cell death primarily through direct action on the cellular function or inhibit or interfere with cell myosis, including alkylating agents, tumour necrosis factors, intercalators, microtubulin inhibitors and topoisomerase inhibitors.

Examples of cytotoxic agents include, but are not limited to, tirapazimine, sertenef, cachectin, ifosfamide, tasonermin, Isonidamine, carboplatin, altretamine, prednimustine, dibromodulcitol, ranimustine, fotemustine, nedaplatin, oxaliplatin, temozolamide, heptaplatin, estramustine, imrosulfan tosylate, trofosfamide, nimustine, dibrospidium chloride, pumitepa, lobaplatin, satraplatin, profiromycin, cisplatin, irofulven, dexifosfamide, cis-aminodichloro(2-methylpyridine)platinum, benzylguanine, glufosfamide, GPX100, (trans,trans,trans)bis-mu-(hexane-1,6-diamine)mu-[diamine-platinum(II)]bis[diamine(chloro)platinum(II)] tetrachloride, diarizidinyl-spermine, arsenic trioxide, 1-(11-dodecylamino-10-hydroxyundecyl)-3,7-dimethylxanthine, zorubicin, idarubicin, daunorubicin, bisantrene, mitoxantrone, pirarubicin, pinafide, valrubicin, amrubicin, antineoplastone, 3'-de-amino-3'-morpholino-13-deoxy-10-hydroxycarminomycin, annamycin, galarubicin, elinafide, MEN10755 and 4-demethoxy-3-deamino-3-aziridinyl-4-methylsulfonyldaunorubicin (see WO 00/50032).

Examples of microtubulin inhibitors include paclitaxel, vindesine sulfate, 3',4'-didehydro-4'-deoxy-8'-norvincaleukoblastine, docetaxol, rhizoxin, dolastatin, mivobulin isethionate, auristatin, cemadotin, RPR109881, BMS184476, vinflunine, cryptophycin, 2,3,4,5,6-pentafluoro-N-(3-fluoro-4-

methoxyphenyl)benzenesulfonamide, anhydrovinblastine, N,N-dimethyl-L-valyl-L-valyl-N-methyl-L-valyl-L-prolyl-L-proline-t-butylamide, TDX258 and BMS188797.

Some examples of topoisomerase inhibitors are topotecan, hycaptamine, irinotecan, rubitecan, 6-ethoxypropionyl-3',4'-O-exobenzylidenechartreusin, 9-methoxy-N,N-dimethyl-5-nitropyrazolo[3,4,5-kl]acridine-2- (6H)propan-amine, 1-amino-9-ethyl-5-fluoro-2,3-dihydro-9-hydroxy-4-methyl-1H,12H-benzo[de]pyrano[3',4':b,7]indolizino[1,2b]quinoline-10,13(9H,15H)dione, lurtotecan, 7-[2-(N-isopropylamino)ethyl]-(20S)camptothecin, BNP1350, BNPI1100, BN80915, BN80942, etoposide phosphate, teniposide, sobuzoxane, 2'-dimethylamino-2'-deoxyetoposide, GL331, N-[2-(dimethylamino)-ethyl]-9-hydroxy-5,6-dimethyl-6H-pyrido[4,3-b]carbazole-1-carboxamide, asulacrine, (5a,5aB,8aa,9b)-9-[2-[N-[2-(dimethylamino)ethyl]-N-methylamino]ethyl]-5-[4-hydroxy-3,5-dimethoxyphenyl]-5,5a,6,8,8a,9-hexo-hydrofuro(3',4':6,7)naphtho(2,3-d)-1,3-dioxol-6-one, 2,3-(methylenedioxy)-5-methyl-7-hydroxy-8-methoxybenzo[c]phenanthridinium, 6,9-bis[(2-aminoethyl)amino]benzo[g]isoquinoline-5,10-dione, 5-(3-aminopropylamino)-7,10-dihydroxy-2-(2-hydroxyethylaminomethyl)-6H-pyrazolo[4,5,1-de]acridin-6-one, N-[1-[2(diethylamino)ethylamino]-7-methoxy-9-oxo-9H-thioxanthene-4-ylmethyl]formamide, N-(2-(dimethylamino)ethyl)acridine-4-carboxamide, 6-[[2-(dimethylamino)ethyl]amino]-3-hydroxy-7H-indeno[2,1-c]quinolin-7-one and dimesna.

"Antiproliferative agents" include antisense RNA and DNA oligonucleotides such as G3139, ODN698, RVASKRAS, GEM231 and INX3001 and anti-metabolites such as enocitabine, carmofur, tegafur, pentostatin, doxifluridine, trimetrexate, fludarabine, capecitabine, galocitabine, cytarabine ocfosfate, fosteabine sodium hydrate, raltitrexed, paltitrexid, emitefur, tiazofurin, decitabine, nolatrexed, pemetrexed, nelzarabine, 2'-deoxy-2'-methylideneцитidine, 2'-fluoromethylene-2'-deoxycytidine, N-[5-(2,3-dihydro-benzofuryl)sulfonyl]-N'-(3,4-dichlorophenyl)urea, N6-[4-deoxy-4-[N2-[2(E),4(E)-tetradecadienoyl]glycylamino]-L-glycero-B-L-mannohepto-pyranosyl]adenine, aplidine, ecteinascidin, troxacicabine, 4-[2-amino-4-oxo-

4,6,7,8-tetrahydro-3H-pyrimidino[5,4-b]-1,4-thiazin-6-yl-(S)-ethyl]-2,5-thienoyl-L-glutamic acid, aminopterin, 5-fluorouracil, alanosine, 11-acetyl-8-(carbamoyloxymethyl)-4-formyl-6-methoxy-14-oxa-1,11-diazatetracyclo-(7.4.1.0.0)tetradeca-2,4,6-trien-9-ylacetic acid ester, swainsonine, lomextroxol, dextrazoxane, methioninase, 2'-cyano-2'-deoxy-N4-palmitoyl-1-B-D-arabinofuranosyl cytosine and 3-aminopyridine-2-carboxaldehyde thiosemicarbazone. "Antiproliferative agents" also include monoclonal antibodies to growth factors other than those listed under "angiogenesis inhibitors", such as trastuzumab, and tumour suppressor genes, such as p53, which can be delivered via recombinant virus-mediated gene transfer (see US Patent No. 6,069,134, for example).

Particular preference is given to the use of the compound according to the invention for the treatment and prophylaxis of tumour diseases.

The tumour is preferably selected from the group of tumours of the squamous epithelium, of the bladder, of the stomach, of the kidneys, of head and neck, of the oesophagus, of the cervix, of the thyroid, of the intestine, of the liver, of the brain, of the prostate, of the urogenital tract, of the lymphatic system, of the stomach, of the larynx and/or of the lung.

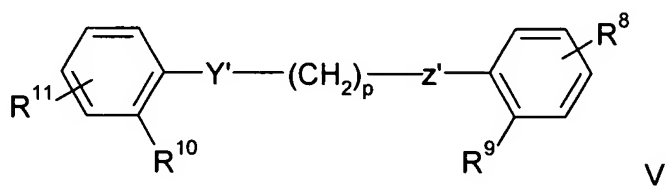
The tumour is furthermore preferably selected from the group lung adenocarcinoma, small-cell lung carcinomas, pancreatic cancer, glioblastomas, colon carcinoma and breast carcinoma.

Preference is furthermore given to the use for the treatment of a tumour of the blood and immune system, preferably for the treatment of a tumour selected from the group of acute myelotic leukaemia, chronic myelotic leukaemia, acute lymphatic leukaemia and/or chronic lymphatic leukaemia.

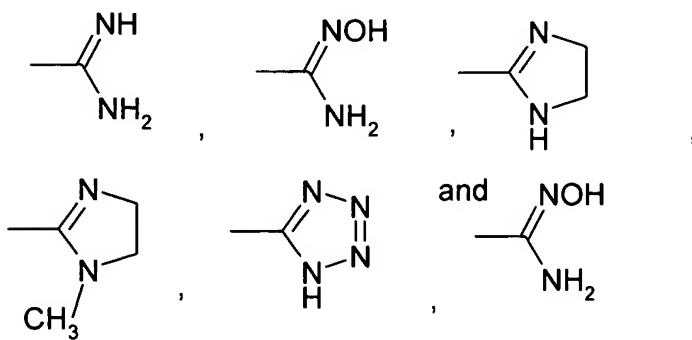
The invention also encompasses a method for the treatment of a patient who has a neoplasm, such as a cancer, by administration of

a) one or more of the compounds of the formula I:

b) and one or more of the compounds of the formula V or acid-addition salts thereof, in particular hydrochlorides:



in which Y' and Z' each, independently of one another, denote O or N, R⁶ and R⁷ each, independently of one another, denote H, OH, halogen, OC1-10-alkyl, OCF₃, NO₂ or NH₂, n denotes an integer between 2 and 6 inclusive, and R⁸ and R⁹ are each, independently of one another, preferably in the meta- or para-position and are selected from the group:



where the first and second compound are administered simultaneously or within 14 days of one another in amounts which are sufficient to inhibit the growth of the neoplasm.

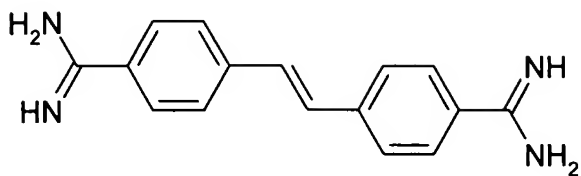
The combination of the compounds of the formula I with the compounds of the formula V and other pentamidine analogues results in a synergistic action in the inhibition of neoplasias. Combinations comprising the compounds of the formula V are mentioned, for example, in WO 02058684.

The mechanism of action of pentamidine or derivatives thereof has not been clearly explained at present: pentamidine or derivatives thereof appears to have pleiotropic actions since it results in a decrease in DNA, RNA and protein synthesis. It was recently described that pentamidine is a capable inhibitor of PRL1, -2 and 3 phosphatases (Pathak et al., 2002) and tyrosine phosphatases, and overexpression thereof is associated with neoplastic malignant tumours in humans. On the other hand, it has been described that pentamidine is a medicament which binds to the DNA minor groove (Puckowska et al., 2004) and is able to exert its action via disturbance of gene expression and/or DNA synthesis.

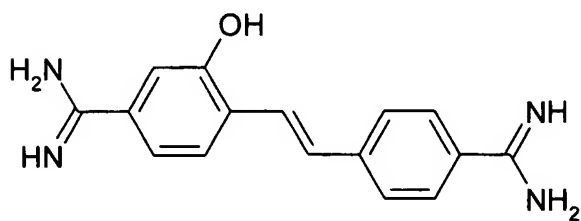
The attached experiments show that:

- both pentamidine and also the compounds of the formula I maintain cells in the G2/M cell cycle.
- the combination of pentamidine and compounds of the formula I have additive to synergistic actions on cell proliferation.

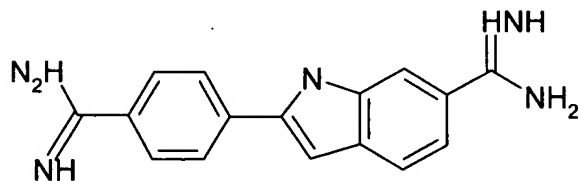
Other suitable pentamidine analogues include stilbamidine (G-1) and hydroxystilbamidine (G-2) and indole analogues thereof (for example G-3):



(G-1)



(G-2) and



(G-3)

Each amidine unit may be replaced, independently of one another, by one of the units defined above for R^8 and R^{11} . As in the case of benzimidazoles and pentamidines, salts of stilbamidine, hydroxystilbamidine and indole derivatives thereof are also suitable for the process according to the invention. Preferred salts include, for example, dihydrochloride and methanesulfonate salts.

Still other analogues are those which fall under a formula which are provided in one of the US Patents No. 5,428,051, 5,521,189, 5,602,172, 5,643,935, 5,723,495, 5,843,980, 6,172,104 and 6,326,395 or the US patent application with the publication No. US 2002/0019437 A1, each of which is incorporated in its entirety by way of reference. Illustrative ana-

logues include 1,5-bis(4'-(N-hydroxyamidino)phenoxy)pentane, 1,3-bis(4'-(N-hydroxyamidino)phenoxy)propane, 1,3-bis(2'-methoxy-4'-(N-hydroxyamidino)phenoxy)propane, 1,4-bis(4'-(N-hydroxyamidino)phenoxy)butane, 1,5-bis(4'-(N-hydroxyamidino)phenoxy)pentane, 1,4-bis(4'-(N-hydroxyamidino)phenoxy)butane, 1,3-bis(4'-(4-hydroxyamidino)phenoxy)propane, 1,3-bis(2'-methoxy-4'-(N-hydroxyamidino)phenoxy)propane, 2,5-bis[4-amidinophenyl]furan, 2,5-bis[4-amidinophenyl]furan bisamide oxime, 2,5-bis[4-amidinophenyl]furan bis-O-methylamide oxime, 2,5-bis[4-amidinophenyl]furan bis-O-ethylamide oxime, 2,8-diamidinodibenzothiophene, 2,8-bis(N-isopropylamidino)carbazole, 2,8-bis(N-hydroxyamidino)carbazole, 2,8-bis(2-imidazolinyl)dibenzothiophene, 2,8-bis(2-imidazolinyl)-5,5-dioxodibenzothiophene, 3,7-diamidinodibenzothiophene, 3,7-bis(N-isopropylamidino)dibenzothiophene, 3,7-diaminodibenzothiophene, 3,7-dibromodibenzothiophene, 3,7-dicyanodibenzothiophene, 2,8-diamidinodibenzofuran, 2,8-di-(2-imidazolinyl)-dibenzofuran, 2,8-di-(N-isopropylamidino)dibenzofuran, 3,7-di-(2-imidazolinyl)dibenzofuran, 3,7-di-(isopropylamidino)dibenzofuran, 3,7-di-(A-hydroxylamidino)dibenzofuran, 2,8-dicyanodibenzofuran, 4,4'-dibromo-2,2'-dinitrobiphenyl, 2-methoxy-2'-nitro-4,4'-dibromobiphenyl, 2-methoxy-2'-amino-4,4'-dibromobiphenyl, 3,7-dibromodibenzofuran, 3,7-dicyanodibenzofuran, 2,5-bis(5-amidino-2-benzimidazolyl)pyrrole, 2,5-bis[5-(2-imidazolinyl)-2-benzimidazolyl]pyrrole, 2,6-bis[5-(2-imidazolinyl)-2-benzimidazolyl]pyridine, 1-methyl-2,5-bis(5-amidino-2-benzimidazolyl)pyrrole, 1-methyl-2,5-bis[5-(2-imidazolinyl)-2-benzimidazolyl]pyrrole, 1-methyl-2,5-bis[5-(1,4,5,6-tetrahydro-2-pyrimidinyl)-2-benzimidazolyl]pyrrole, 2,6-bis(5-amidino-2-benzimidazolyl)pyridine, 2,6-bis[5-(1,4,5,6-tetrahydro-2-pyrimidinyl)-2-benzimidazolyl]pyridine, 2,5-bis(5-amidino-2-benzimidazolyl)furan, 2,5-bis[5-(2-imidazolinyl)-2-benzimidazolyl]furan, 2,5-bis(4-guanylphenyl)furan, 2,5-bis(4-guanylphenyl)-3,4-dimethylfuran, 2,5-di-p-[2-(3,4,5,6-tetrahydropyrimidinyl)phenyl]furan, 2,5-bis[4-(2-imidazolinyl)phenyl]furan, 2,5-[bis{4-(2-tetrahydropyrimidinyl)}phenyl]p-(tolyloxy)furan, 2,5-[bis-

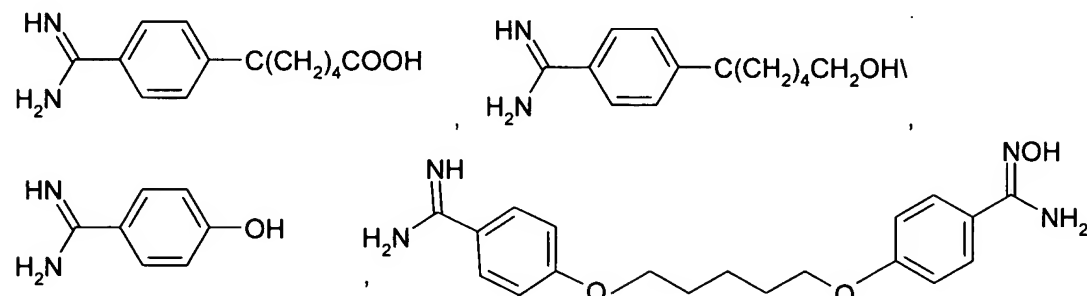
{4-(2-imidazolinyl)phenyl]-3-p-(tolyloxy)furan, 2,5-bis{4-[5-(N-2-aminoethylamido)benzimidazol-2-yl]phenyl}furan, 2,5-bis[4-(3a,4,5,6,7,7a-hexahydro-1H-benzimidazol-2-yl)phenyl]furan, 2,5-bis[4-(4,5,6,7-tetrahydro-1H-1,3-diazepin-2-yl)phenyl]furan, 2,5-bis(4-N,N-dimethylcarboxyhydrazidophenyl)furan, 2,5-bis{4-[2-(N-2-hydroxyethyl)imidazolinyl]phenyl}furan, 2,5-bis[4-(N-isopropylamidino)phenyl]furan, 2,5-bis{4-[3-(dimethylaminopropyl)amidino]phenyl}furan, 2,5-bis{4-[N-(3-aminopropyl)amidino]phenyl}furan, 2,5-bis[2-(imidazolinyl)phenyl]-3,4-bis(methoxymethyl)furan, 2,5-bis[4-N-(dimethylaminoethyl)guanyl]phenylfuran, 2,5-bis{4-[(N-2-hydroxyethyl)guanyl]phenyl}furan, 2,5-bis[4-N-(cyclopropylguanyl)phenyl]furan, 2,5-bis[4-(N,N-diethylaminopropyl)guanyl]phenylfuran, 2,5-bis{4-(N-ethylimidazolinyl)phenyl}furan, 2,5-bis{4-[N-(3-pentylguanyl)]phenyl}furan, 2,5-bis[4-(2-imidazolinyl)phenyl]-3-methoxyfuran, 2,5-bis[4-(N-isopropylamidino)phenyl]-3-methylfuran, bis[5-amidino-2-benzimidazoly]-methane, bis[5-(2-imidazoly)-2-benzimidazoly]methane, 1,2-bis[5-amidino-2-benzimidazoly]ethane, 1,2-bis[5-(2-imidazoly)-2-benzimidazoly]ethane, 1,3-bis[5-amidino-2-benzimidazoly]propane, 1,3-bis[5-(2-imidazoly)-2-benzimidazoly]propane, 1,4-bis[5-(2-imidazoly)-2-benzimidazoly]butane, 1,8-bis[5-amidino-2-benzimidazoly]octane, trans-1,2-bis[5-amidino-2-benzimidazoly]ethene, 1,4-bis[5-(2-imidazoly)-2-benzimidazoly]-1-butene, 1,4-bis[5-(2-imidazoly)-2-benzimidazoly]-2-butene, 1,4-bis[5-(2-imidazoly)-2-benzimidazoly]-1-methylbutane, 1,4-bis[5-(2-imidazoly)-2-benzimidazoly]-2-ethylbutane, 1,4-bis[5-(2-imidazoly)-2-benzimidazoly]-1-methyl-1-butene, 1,4-bis[5-(2-imidazoly)-2-benzimidazoly]-2,3-diethyl-2-butene, 1,4-bis[5-(2-imidazoly)-2-benzimidazoly]-1,3-butadiene, 1,4-bis[5-(2-imidazoly)-2-benzimidazoly]-2-methyl-1,3-butadiene, bis[5-(2-pyrimidyl)-2-benzimidazoly]methane, 1,2-bis[5-(2-pyrimidyl)-2-benzimidazoly]ethane, 1,3-bis[5-amidino-2-benzimidazoly]propane, 1,3-bis[5-(2-pyrimidyl)-2-benzimidazoly]propane, 1,4-bis[5-(2-pyrimidyl)-2-benzimidazoly]-1-butene, 1,4-bis[5-(2-pyrimidyl)-2-benzimidazoly]-2-butene, 1,4-bis[5-(2-pyrimidyl)-2-benzimidazoly]-1-methylbutane, 1,4-bis[5-(2-pyrimidyl)-2-benzimidazoly]

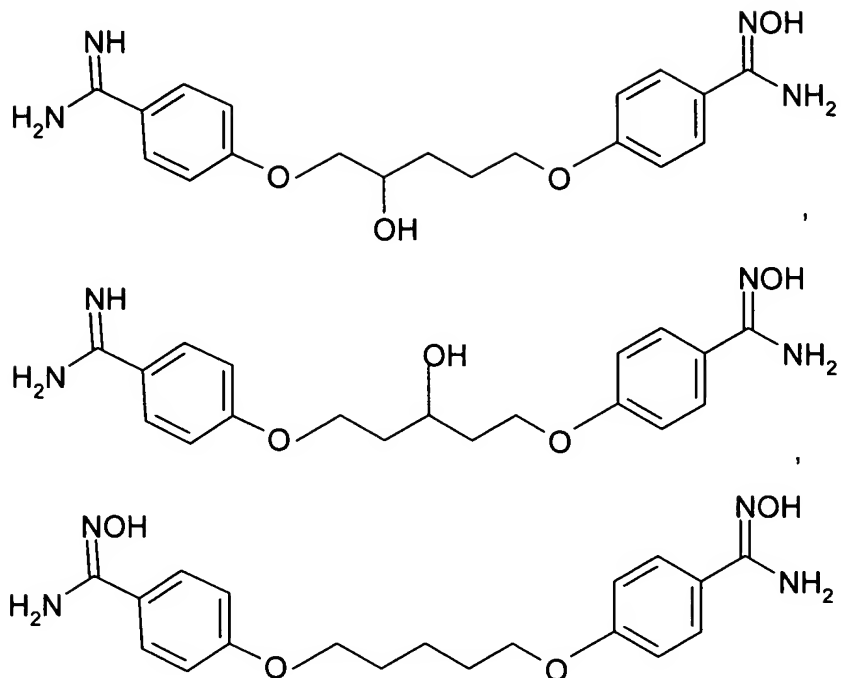
benzimidazolyl]-2-ethylbutane, 1,4-bis[5-(2-pyrimidyl)-2-benzimidazolyl]-1-methyl-1-butene, 1,4-bis[5-(2-pyrimidyl)-2-benzimidazolyl]-2,3-diethyl-2-butene, 1,4-bis[5-(2-pyrimidyl)-2-benzimidazolyl]-1,3-butadiene and 1,4-bis[5-(2-pyrimidyl)-2-benzimidazolyl]-2-methyl-1,3-butadiene, 2,4-bis(4-guanylphenyl)pyrimidine, 2,4-bis(4-imidazolin-2-yl)pyrimidine, 2,4-bis-[(tetrahydropyrimidinyl-2-yl)phenyl]pyrimidine, 2-(4-[N-i-propylguanyl]-phenyl)-4-(2-methoxy-4-[N-i-propylguanyl]phenyl)pyrimidine, 4-(N-cyclopentylamidino)-1,2-phenylenediamine, 2,5-bis[2-(5-amidino)benzimidazoyl]-furan, 2,5-bis[2-{5-(2-imidazolino)}benzimidazoyl]furan, 2,5-bis[2-(5-N-isopropylamidino)benzimidazoyl]furan, 2,5-bis[2-(5-N-cyclopentylamidino)-benzimidazoyl]furan, 2,5-bis[2-(5-amidino)benzimidazoyl]pyrrole, 2,5-bis[2-{5-(2-imidazolino)}benzimidazoyl]pyrrole, 2,5-bis[2-(5-N-isopropylamidino)-benzimidazoyl]pyrrole, 2,5-bis[2-(5-N-cyclopentylamidino)-benzimidazoyl]pyrrole, 1-methyl-2,5-bis[2-(5-amidino)benzimidazoyl]pyrrole, 2,5-bis[2-{5-(2-imidazolino)}benzimidazoyl]-1-methylpyrrole, 2,5-bis[2-(5-N-cyclopentylamidino)benzimidazoyl]-1-methylpyrrole, 2,5-bis[2-(5-N-isopropylamidino)benzimidazoyl]thiophene, 2,6-bis[2-{5-(2-imidazolino)}-benzimidazoyl]pyridine, 2,6-bis[2-(5-amidino)benzimidazoyl]pyridine, 4,4'-bis[2-(5-N-isopropylamidino)benzimidazoyl]-1,2-diphenylethane, 4,4'-bis[2-(5-N-cyclopentylamidino)benzimidazoyl]-2,5-diphenylfuran, 2,5-bis[2-(5-amidino)benzimidazoyl]benzo[b]furan, 2,5-bis[2-(5-N-cyclopentylamidino)-benzimidazoyl]benzo[b]furan, 2,7-bis[2-(5-N-isopropylamidino)benzimidazoyl]fluorine, 2,5-bis[4-(3-(N-morpholinopropyl)carbamoyl)phenyl]furan, 2,5-bis[4-(2-N,N-dimethylaminoethylcarbamoyl)phenyl]furan, 2,5-bis[4-(3-N,N-dimethylaminopropylcarbamoyl)phenyl]furan, 2,5-bis[4-(3-N-methyl-3-N-phenylaminopropylcarbamoyl)phenyl]furan, 2,5-bis[4-(3-N,N8,N11-trimethylaminopropylcarbamoyl)phenyl]furan, 2,5-bis[3-amidinophenyl]-furan, 2,5-bis[3-(N-isopropylamidino)amidinophenyl]furan, 2,5-bis[3-[(N-(2-dimethylaminoethyl)amidino)phenyl]furan, 2,5-bis[4-(N-2,2,2-trichloroethoxy-carbonyl)amidinophenyl]furan, 2,5-bis[4-(N-thioethylcarbonyl)amidinophenyl]furan, 2,5-bis[4-(N-benzylloxycarbonyl)amidinophenyl]furan, 2,5-bis[4-(N-phenoxy carbonyl)amidinophenyl]furan, 2,5-bis[4-(N-(4-fluoro)-

phenoxy carbonyl)amidinophenyl]furan, 2,5-bis[4-(N-(4-methoxy)phenoxy carbonyl)amidinophenyl]furan, 2,5-bis[4-(1-acetoxyethoxycarbonyl)amidinophenyl]furan and 2,5-bis[4-(N-(3-fluoro)phenoxy carbonyl)amidinophenyl]furan. Processes for the preparation of one of the above compounds are described in US patents No. 5,428,051, 5,521,189, 5,602,172, 5,643,935, 5,723,495, 5,843,980, 6,172,104 and 6,326,395 or the US patent application with the publication No. US 2002/0019437 A1.

Pentamidine metabolites are likewise suitable in the antiproliferative combination according to the invention. Pentamidine is rapidly metabolised in the body to at least seven primary metabolites. Some of these metabolites have one or more actions in common with pentamidine. Pentamidine metabolites have an antiproliferative action when combined with a benzimidazole or an analogue thereof.

Seven pentamidine analogues are shown below.





The combinations according to the invention of compounds of the formula I and formula V or analogues thereof and metabolites thereof are suitable for the treatment of neoplasms. Combination therapy can be carried out alone or in combination with another therapy (for example operation, irradiation, chemotherapy, biological therapy). In addition, a person whose risk of developing a neoplasm is greater (for example someone who is genetically predisposed or someone who previously had a neoplasm) can be given prophylactic treatment in order to inhibit or delay neoplasm formation.

The invention likewise relates to the combination of kinesin ATPase Eg5/KSP with the compounds of the formula V, pentamidine, analogues thereof and/or metabolites thereof.

The dosage and frequency of administration of each compound in the combination can be controlled independently. For example, one compound may be administered orally three times daily, while the second compound may be administered intramuscularly once per day. The compounds may also be formulated together, leading to administration of both compounds.

The antiproliferative combinations according to the invention can also be provided as components of a pharmaceutical package. The two medicaments can be formulated together or separately and in individual dosage amounts.

In another aspect, the invention encompasses a method for the treatment of a patient who has a neoplasm, such as a cancer, by administration of a compound of the formula (I) and (V) in combination with an antiproliferative agent. Suitable antiproliferative agents encompass those provided in Table 1.

Above and below, all temperatures are indicated in °C. In the following examples, "conventional work-up" means: if necessary, water is added, the pH is adjusted, if necessary, to values between 2 and 10, depending on the constitution of the end product, the mixture is extracted with ethyl acetate or dichloromethane, the phases are separated, the organic phase is dried over sodium sulfate and evaporated, and the product is purified by chromatography on silica gel and/or by crystallisation. Rf values on silica gel; eluent: ethyl acetate/methanol 9:1.

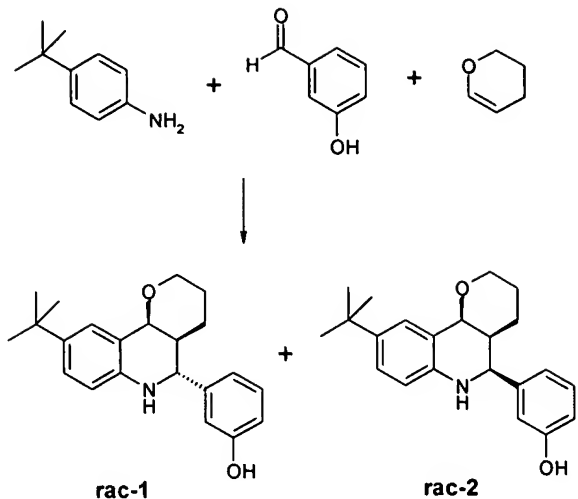
Mass spectrometry (MS): EI (electron impact ionisation) M⁺

FAB (fast atom bombardment) (M+H)⁺

ESI (electrospray ionisation) (M+H)⁺

APCI-MS (atmospheric pressure chemical ionisation – mass spectrometry)
(M+H)⁺

Example 1



a. Reaction in the presence of trifluoroacetic acid (TFA)

The solution of the TFA salt of 4-*tert*-butylaniline in acetonitrile (4-*tert*-butylaniline (0.50 g, 3.35 mmol) was taken up in acetonitrile (4 ml), cooled to 0°C, and TFA (0.38 g, 3.35 mmol) was slowly added with stirring) was added rapidly to a solution, cooled to 0°C, of 3-hydroxybenzaldehyde (0.41 g, 3.35 mmol) and 3,4-dihydro-2H-pyran (0.28 g, 3.35 mmol) in acetonitrile (2 ml), and the mixture was stirred at this temperature for a further 60 min. The reaction solution was evaporated to dryness and separated by column chromatography, giving a colourless solid (620 mg, 1.84 mmol, 55%), which proved to be an isomer mixture of the respective racemic *trans/cis* compound in the ratio 13:1.

In order to separate the *cis/trans* isomers, the solid was taken up in 0.1 N HCl in isopropanol (20 ml), 100 ml of each of diethyl ether and cyclohexane were added, and the mixture was crystallised overnight at 4°C. The colourless solid was filtered off, washed with a little diethyl ether and dried, giving 570 mg (1.52 mmol) of the colourless compound rac-1 hydrochloride. The mother liquor was evaporated to dryness in a rotary evaporator, and

the *cis* isomer was purified by column chromatography and converted into the hydrochloride (70 mg, 0.19 mmol of a colourless solid).

Relatively small amounts of rac-1 and rac-2 were separated into the corresponding enantiomers by chiral HPLC (2x Chiraldak AD 20 µm, eluent: methanol) and again converted into the hydrochlorides.

b. Reaction in the presence of hexafluoroisopropanol (HFIP)

In order to obtain higher yields of the *cis* isomer, hexafluoroisopropanol (HFIP) was employed instead of TFA.

4-*tert*-Butylaniline (0.50 g, 3.35 mmol), 3-hydroxybenzaldehyde (0.41 g, 3.35 mmol) and 3,4-dihydro-2H-pyran (0.28 g, 3.35 mmol) were dissolved in acetonitrile (1 ml), HFIP (0.56 g, 3.35 mmol) was added dropwise at RT, and the mixture was stirred at RT for a further 18 h. The reaction solution was evaporated to dryness and separated by column chromatography, giving a colourless solid (485 mg, 1.44 mmol, 43%), which proved to be an isomer mixture of the respective racemic *trans/cis* compound in the ratio 1:1.6.

Separation of the isomers was carried out as described under a.).

c. Reaction in the presence of bismuth(III) chloride

3-Hydroxybenzaldehyde (0.41 g, 3.35 mmol) and 4-*tert*-butylaniline (0.50 g, 3.35 mmol), each dissolved in acetonitrile (1 ml), were added at RT to a suspension of BiCl₃ (0.21 g, 0.67 mmol) and anhydrous sodium sulfate (0.40 g) in acetonitrile (2 ml), and the mixture was stirred at RT for 10 min. 3,4-Dihydro-2H-pyran (0.28 g, 3.35 mmol) was then added dropwise at RT, and the mixture was stirred at RT for a further 30 min. The reaction solution was filtered through Celite, the filtrate was evaporated to dryness and purified by column chromatography, giving a colourless solid (850 mg,

2.51 mmol, 75%), which proved to be an isomer mixture of the respective racemic *trans/cis* compound in the ratio 1:1.1.

Separation of the isomers was carried out as described under a.) .

d. Reaction in the presence of ytterbium(III) triflate

3-Hydroxybenzaldehyde (0.41 g, 3.35 mmol) and 4-*tert*-butylaniline (0.50 g, 3.35 mmol), each dissolved in acetonitrile (1 ml), were added at RT to a suspension of Yb(OTf)₃ (0.42 g, 0.67 mmol) and anhydrous sodium sulfate (0.40 g) in acetonitrile (2 ml), and the mixture was stirred at RT for 10 min. 3,4-Dihydro-2H-pyran (0.28 g, 3.35 mmol) was then added dropwise at RT, and the mixture was stirred at RT for a further 30 min. The reaction solution was filtered through Celite, the filtrate was evaporated to dryness and purified by column chromatography, giving a colourless solid (780 mg, 2.31 mmol, 69%), which proved to be an isomer mixture of the respective racemic *trans/cis* compound in the ratio 1:1.3.

Separation of the isomers was carried out as described under a.) .

e. Reaction in the presence of scandium(III) triflate

3-Hydroxybenzaldehyde (0.41 g, 3.35 mmol) and 4-*tert*-butylaniline (0.50 g, 3.35 mmol), each dissolved in acetonitrile (1 ml), were added at RT to a suspension of Sc(OTf)₃ (0.33 g, 0.67 mmol) and anhydrous sodium sulfate (0.40 g) in acetonitrile (2 ml), and the mixture was stirred at RT for 10 min. 3,4-Dihydro-2H-pyran (0.28 g, 3.35 mmol) was then added dropwise at RT, and the mixture was stirred at RT for a further 30 min. The reaction solution was filtered through Celite, the filtrate was evaporated to dryness and purified by column chromatography, giving a colourless solid (620 mg, 1.84 mmol, 55%), which proved to be an isomer mixture of the respective racemic *trans/cis* compound.

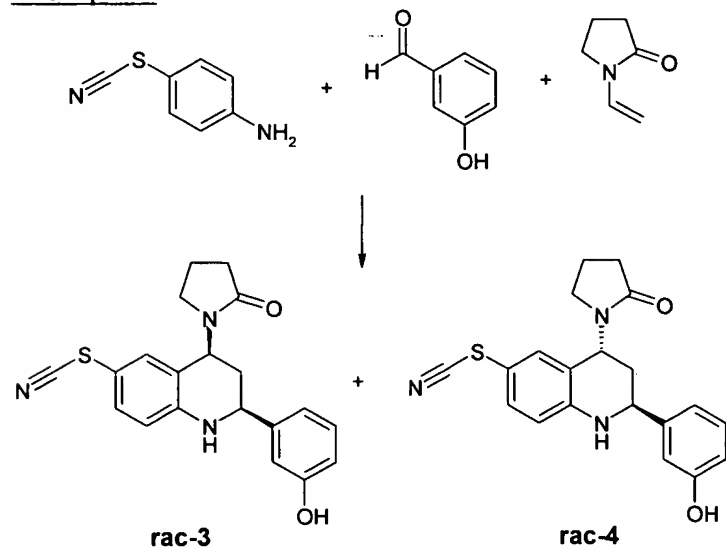
Separation of the isomers was carried out as described under a.) .

f. Reaction in the presence of cerium(IV) ammonium nitrate (CAN)

CAN (0.37 g, 0.67 mmol) was added at RT to a solution of 3-hydroxybenzaldehyde (0.41 g, 3.35 mmol), 4-*tert*-butylaniline (0.50 g, 3.35 mmol) and 3,4-dihydro-2H-pyran (0.28 g, 3.35 mmol) in acetonitrile (10 ml), and the mixture was stirred at RT for a further 30 min. The reaction mixture was evaporated to dryness and purified by column chromatography, giving a colourless solid (237 mg, 0.70 mmol, 21%), which proved to be an isomer mixture of the respective racemic *trans/cis* compound in the ratio 5:1.

Separation of the isomers was carried out as described under a.).

Example 2



a. Reaction in the presence of trifluoroacetic acid (TFA)

The solution of the TFA salt of 4-thiocyanatoaniline in acetonitrile (thiocyanatoaniline (0.46 g, 3.06 mmol) was taken up in acetonitrile (4 ml), cooled to 0°C, and TFA (0.35 g, 3.06 mmol) was slowly added with stirring) was added rapidly to a solution, cooled to 0°C, of 3-hydroxybenz-

aldehyde (0.37 g, 3.06 mmol) and 1-vinyl-2-pyrrolidinone (0.34 g, 3.06 mmol) in acetonitrile (2 ml), and the mixture was stirred at this temperature for a further 60 min and at RT for 18 h. The reaction solution was evaporated, taken up using a little ethyl acetate and crystallised using diethyl ether, giving a colourless solid (390 mg, 1.06 mmol, 35%), which proved to be the *cis* isomer.

In order to separate the *cis/trans* isomers, the filtrate was evaporated to dryness and purified by column chromatography, giving the *trans* isomer as colourless solid (160 mg, 0.44 mmol, 14%).

Relatively small amounts of rac-3 and rac-4 were separated into the corresponding enantiomers by chiral HPLC (2x Chiraldak AD 20 µm, eluent: methanol) and converted into the hydrochlorides.

The following compounds according to the invention are obtained analogously using the corresponding precursors. The retention times indicated were obtained under the following conditions:

Method A:

Merck Hitachi LaChrom

Gradient 9 min

Flow rate : 1.5 ml/min from 80:20 to 0:100 (within 6 min), 0:100 (1 min), back to 80:20 (within 1 min), water/acetonitrile

Water + TFA (0.1% vol)

Acetonitrile + TFA (0.1% vol)

Column : LichroCART 125-4 RP- LiChrospher 60, RP-select-B (5 µm/60 mm)

Wavelength : 220 nm

Method B:

Merck Hitachi LaChrom

Gradient 9 min

Flow rate : 1.5 ml/min from 80:20 to 0:100 (within 6 min), 0:100 (1 min), back to 80:20 (within 0.8 min), water/acetonitrile

Water + TFA (0.1% vol)

Acetonitrile + TFA (0.1% vol)

Column : Lichrospher RP-select-B (5 μ m/125 mm)

Wavelength : 220 nm

Method C:

Agilent 1100 Series

Gradient 3.5 min

Flow rate : 2 ml/min from 80:20 to 0:100 (within 2.3 min), 0:100 (0.5 min),
back to 80:20, water/acetonitrile

Water + TFA (0.1% vol)

Acetonitrile + TFA (0.1% vol)

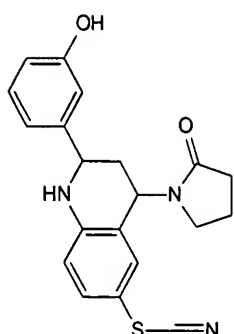
Column : β test research sample, Chromolith Performance RP-18e
(3 μ m/100 mm)

Wavelength : 220 nm

Examples 3 - 654

		[M+1] ⁺	Retention times [min]	Method
(3)		308	5.71	A
(4)		340	6.16 / 6.40	A
(5)		312	5.31	A

(6)

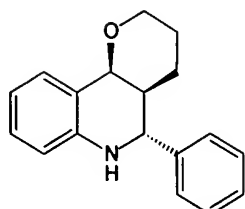


366

4.4

A

(7)

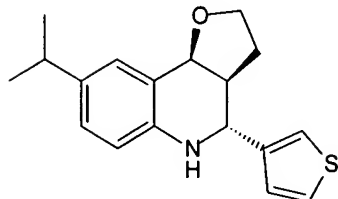


266

5.28

A

(8)



300

5.2

A

(9)

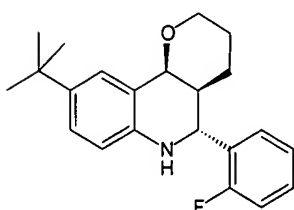


266

5.6

A

(10)

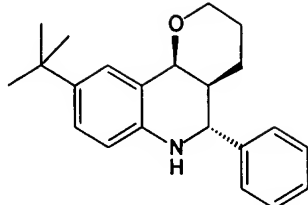


340

6.16

A

(11)

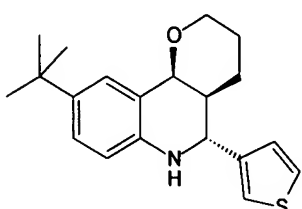


322

5.92

A

(12)



328

5.65

A

(13)

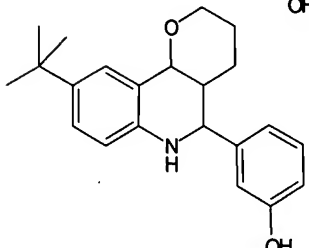


365

4.51

A

(14)

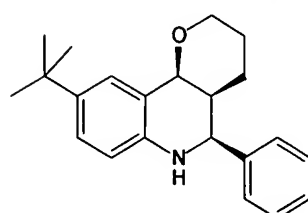


338

5.11

A

(15)

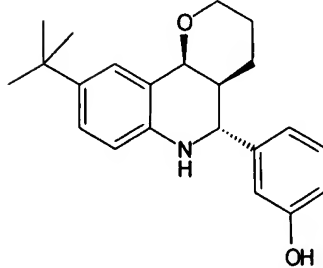


322

6.29

A

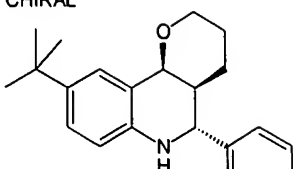
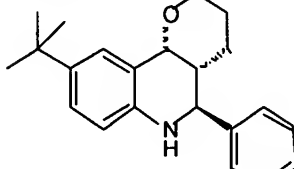
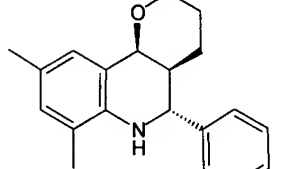
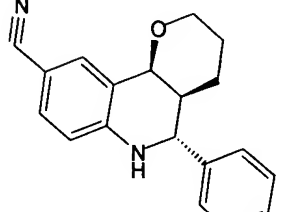
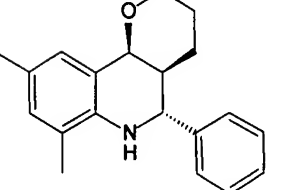
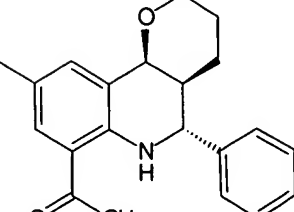
(16)



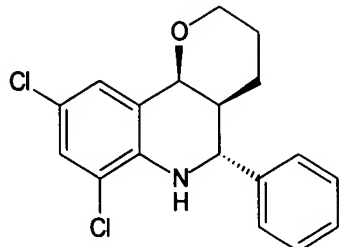
338

5.12

A

(17)	CHIRAL 	338	5.12	A
(18)	CHIRAL 	338	5.12	A
(19)		310	5.08	A
(20)		291	5.31	A
(21)		294	5.68	A
(22)		324	5.36	A

(23)

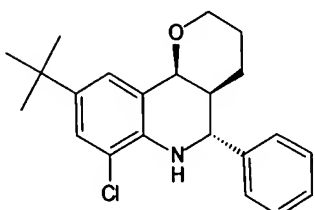


335

6.4

A

(24)

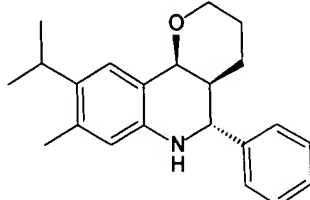


357

3.224

C

(25)

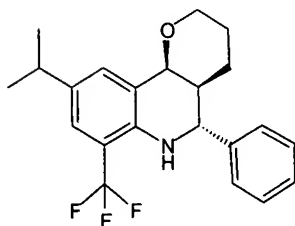


322

2.888

C

(26)

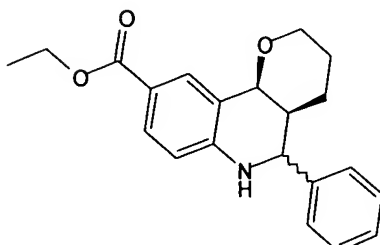


376

6.72

A

(27)

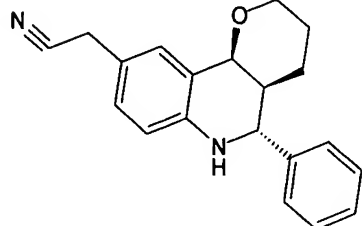


338

5.65/ 5.76

A

(28)

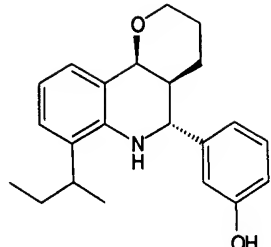


305

5.2

A

(29)

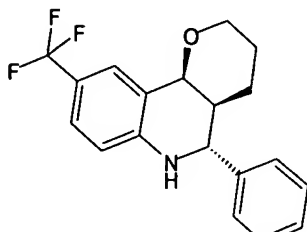


338

5.54

A

(30)

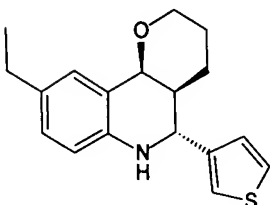


334

5.97

A

(31)

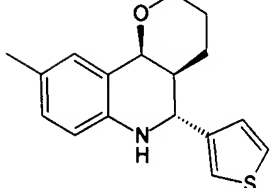


300

5.15

A

(32)

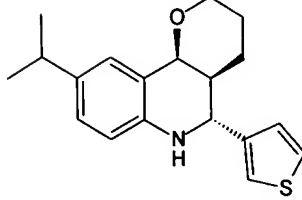


286

4.69

A

(33)

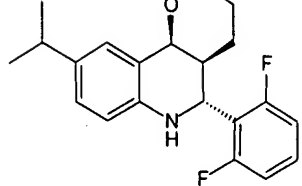


314

5.39

A

(34)

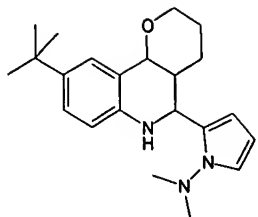


344

6.05

A

(35)

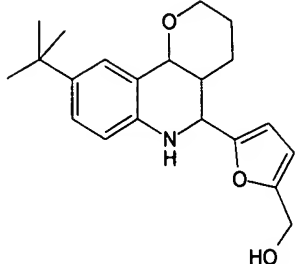


355

5.33

A

(36)

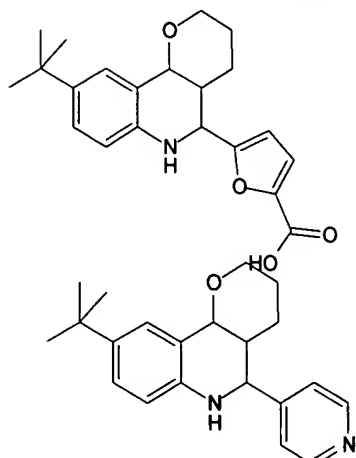


342

4.77

A

(37)



356

5.04

A

(38)

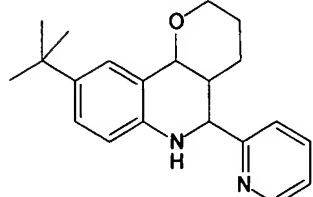


323

4.24

A

(39)

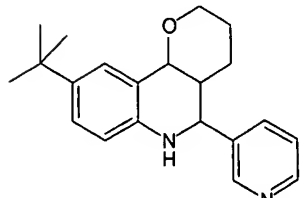


323

4.21

A

(40)



323

4.21

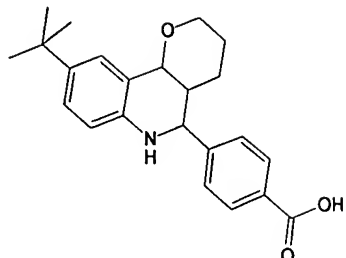
A

(41)		294	5.41	A
(42)		328	6.08	A
(43)		322	6	A
(44)		323	5.68	A
(45)		350	5.52	A
(46)		310	4.61 / 4.67	A

(47)

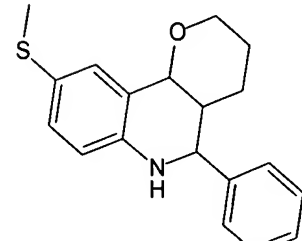
350 6.08 A

(48)



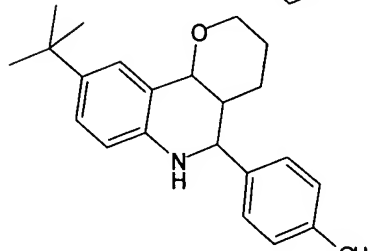
366 5.33/ 5.52 A

(49)



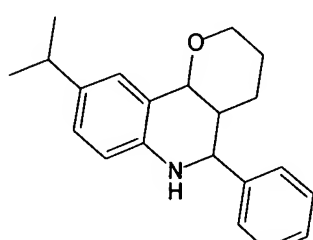
312 4.13/ 5.65 A

(50)



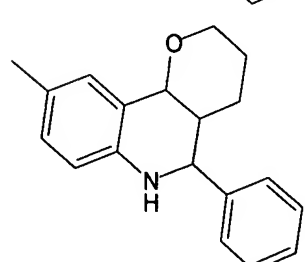
338 4.85 A

(51)

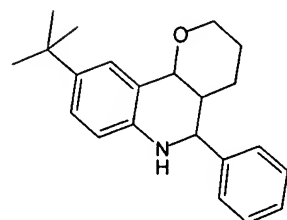


308 5.68 A

(52)



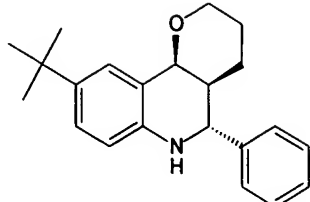
280 5.15 A



(53)

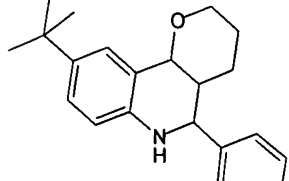
366 5.28 A

(54)



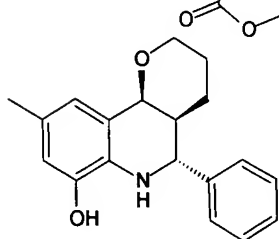
322 5.95 A

(55)



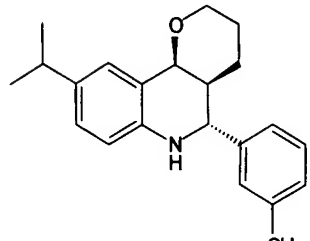
381 6.11 A

(56)



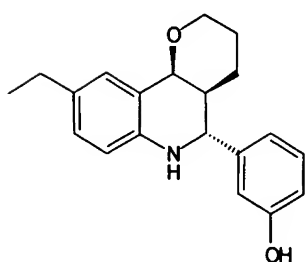
296 4.27 A

(57)

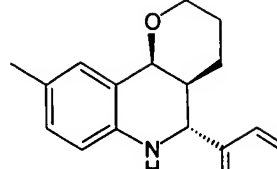
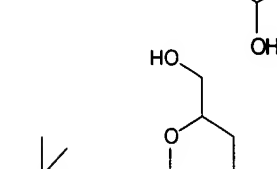
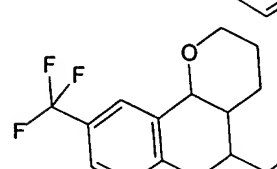
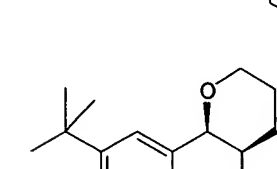
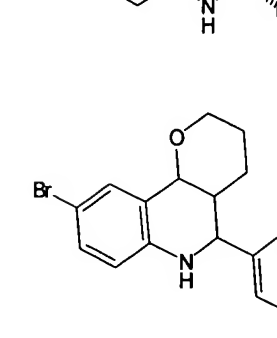
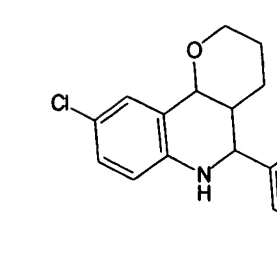


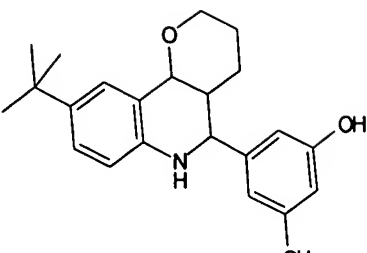
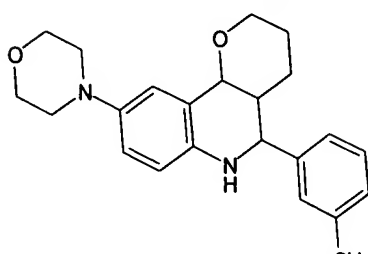
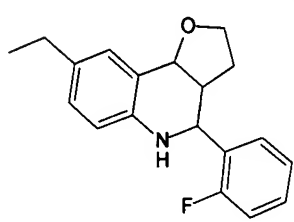
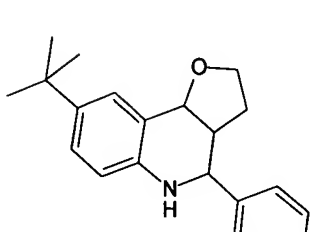
324 4.93 A

(58)



310 2.227 C

(59)		296	4.24	A
(60)		352	5.17 / 5.33	A
(61)		350	5.31	A
(62)		338	5.09	A
(63)		360/362	5.17	A
(64)		317	5.12	A

(65)		354	4.48	A
(66)		367	2.80 ?	A
(67)		298	5.52	A
(68)		284	5.25	A
(69)		324	4.96	A
(70)		367	6.16	A

(71)		382	5.6	A
(72)		300	4.59	A
(73) CHIRAL		322	6.11	A
(74) CHIRAL		322	6.11	A
(75)		322	5.87	A
(76)		338	5.31	A

(77)

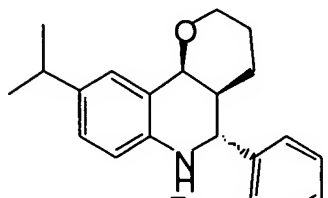


386/388

6.32

A

(78)



376

6.35

A

(79)

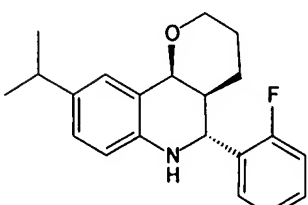


343

6.24

A

(80)

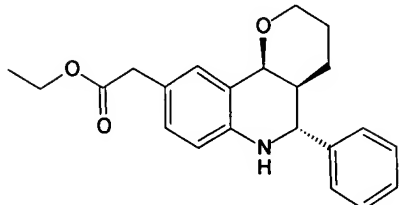


326

6.03

A

(81)

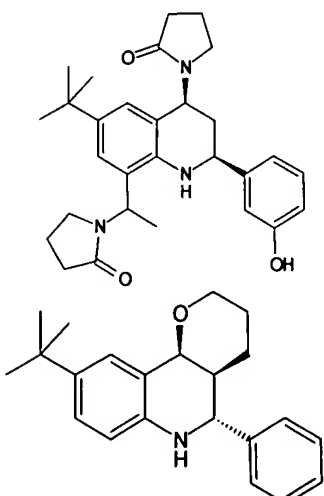


352

5.47

A

(82)



477

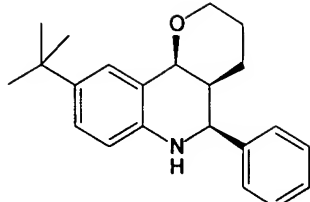
4.8

A

(83)

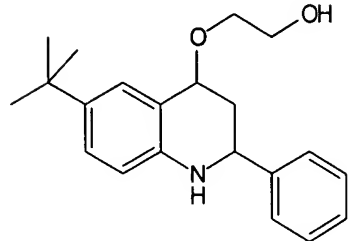
338 5.15 A

(84)



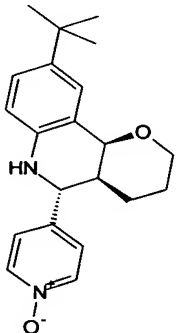
322 6.32 A

(85)



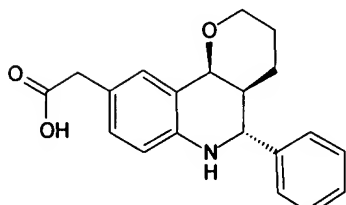
326 5.2 A

(86)



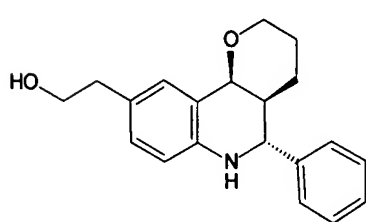
339 4.8 A

(87)



324 4.4 A

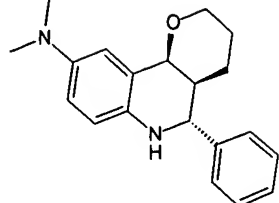
(88)



310 4.16 A

(89)		435	3.95	A
(90)		322	6.11	A
(91)		353	6.13	A
(92) CHIRAL		322	6.45	A
(93) CHIRAL		322	6.45	A
(94)		359	4.27	A

(95)

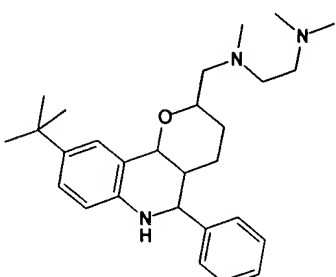


309

3.87

A

(96)

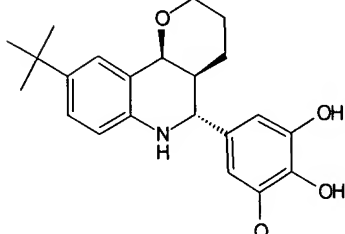


437

4.03 / 4.13

A

(97)

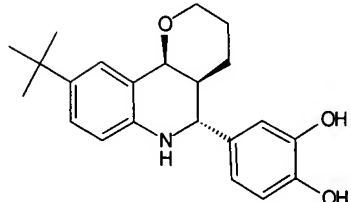


384

4.51

A

(98)

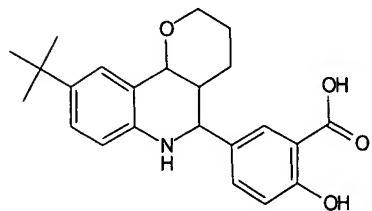


354

4.48

A

(99)

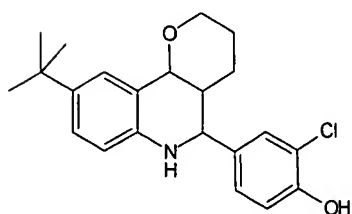


382

5.33

A

(100)

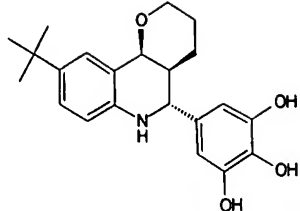


373

5.52

A

(101)

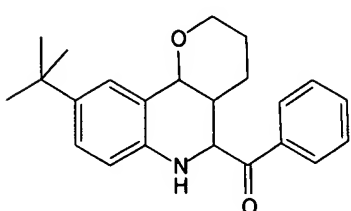


370

4.11

A

(102)

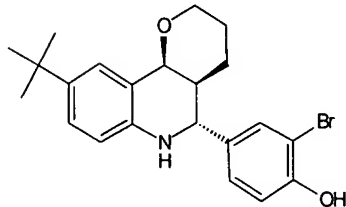


350

6.05

A

(103)

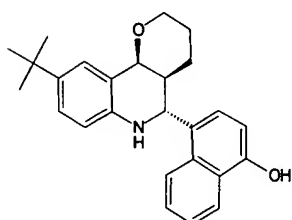


416/418

5.57

A

(104)

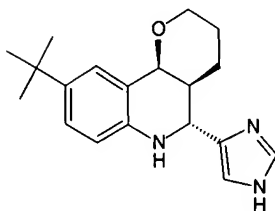


389

5.65

A

(105)

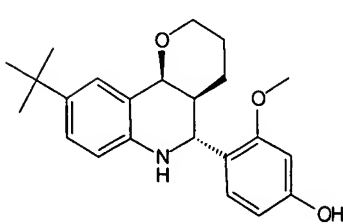


312

3.97

A

(106)

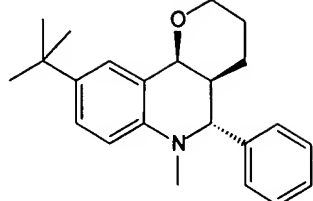


368

4.67

A

(107)

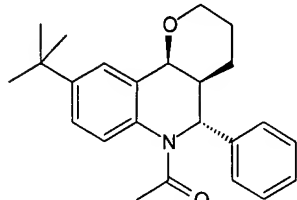


336

3.015

C

(108)

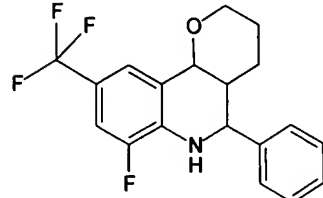


365

2.423

C

(109)

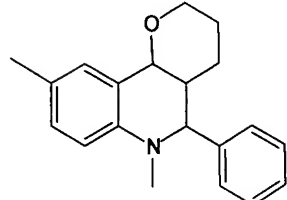


352

6.13

A

(110)

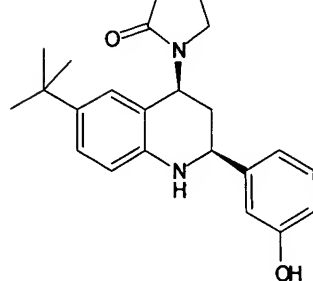


294

2.697

C

(111)

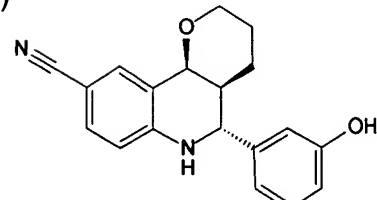


365

4.53

A

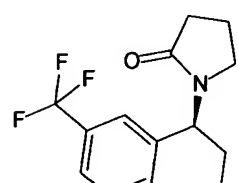
(112)



307

4.61

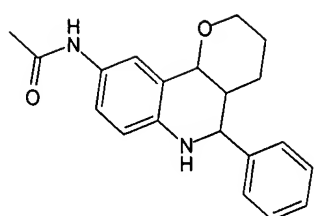
A



(113)

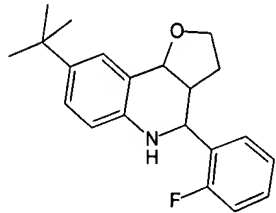
377 4.67 A

(114)



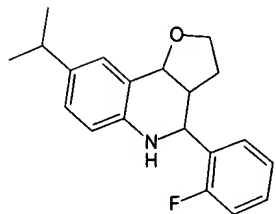
323 4.085/ 4.45 A

(115)



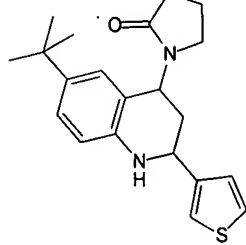
326 5.87 A

(116)



312 2.504 C

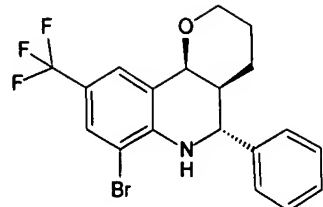
(117)



356 4.91 A

- 100 -

(118)

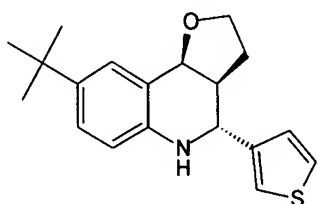


412/414

6.51

A

(119)

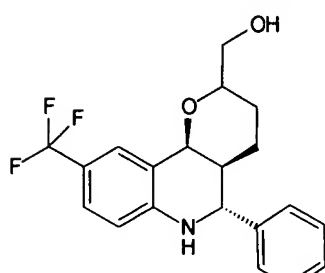


314

5.47

A

(120)

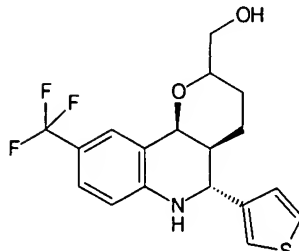


364

5.20 / 5.39

A

(121)

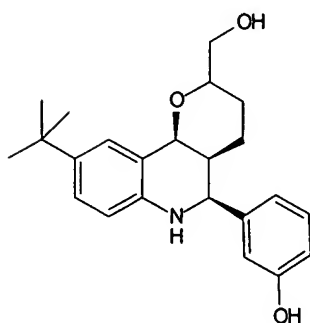


370

5.25

A

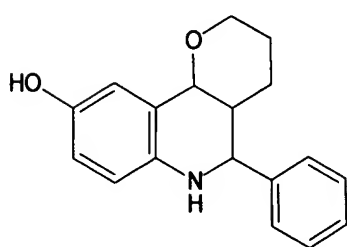
(122)



368

4.69

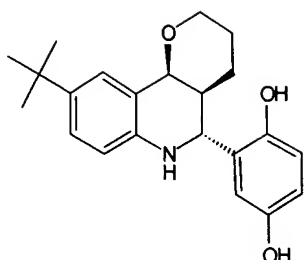
A



(123)

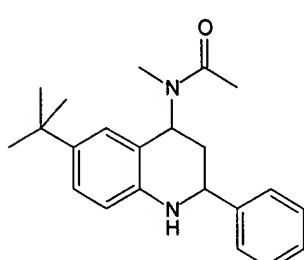
282 3.12 A

(124)



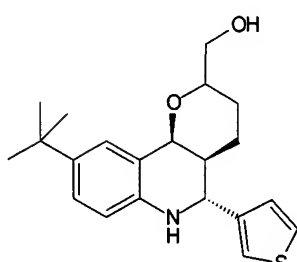
354 4.45 A

(125)



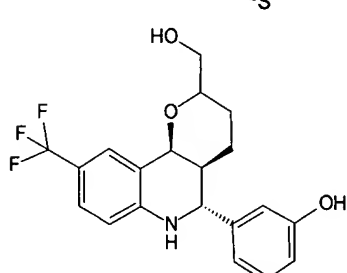
337 4.56 A

(126)



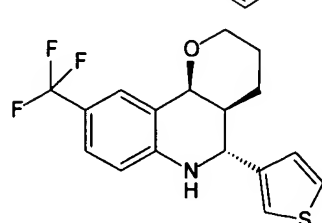
359 4.88 A

(127)



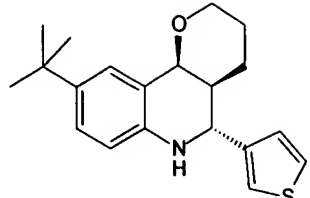
380 4.72 A

(128)



340 5.92 A

(129)

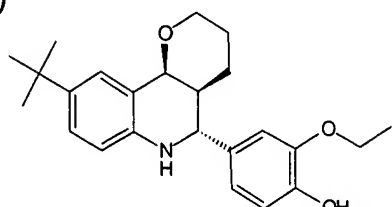


328

5.65

A

(130)

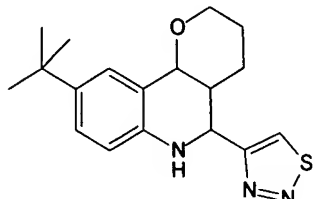


383

5.23

A

(131)

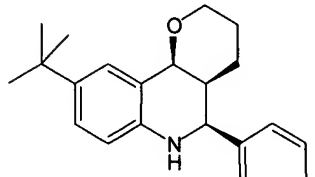


330

5.41 / 5.55

A

(132)

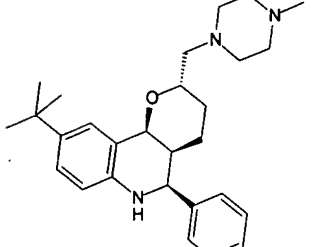


338

5.17

A

(133)

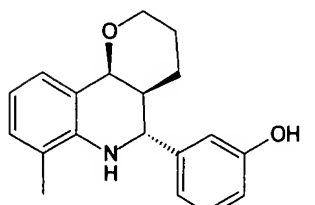


435

3.97

A

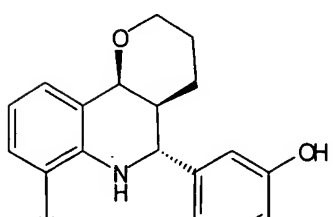
(134)



296

4.83

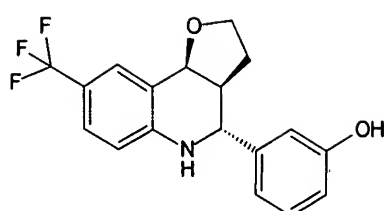
A



(135)

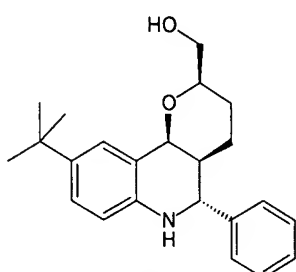
307 4.72 A

(136)



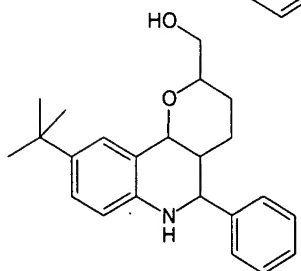
336 5.01 A

(137)



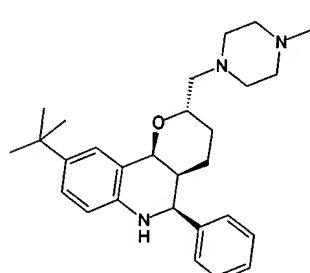
352 5.15 A

(138)



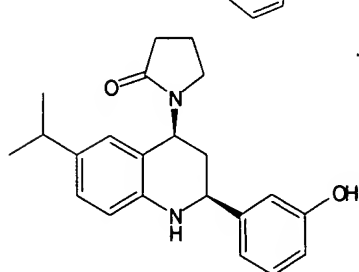
352 5.15 / 5.33 A

(139)

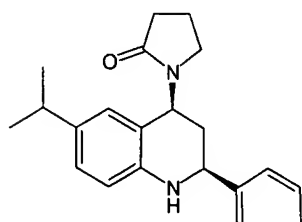


435 3.97 A

(140)



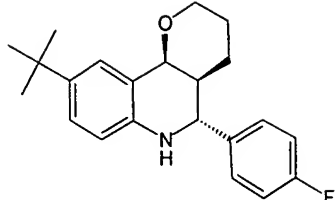
351 4.29 A



(141)

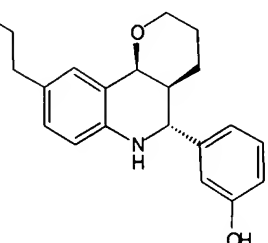
351 3.97 A

(142)



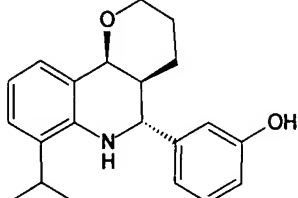
340 2.87 C

(143)



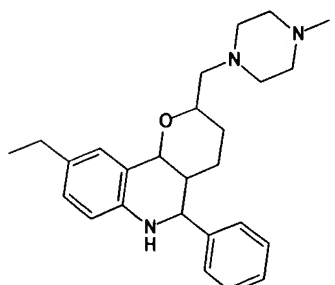
338 2.414 C

(144)



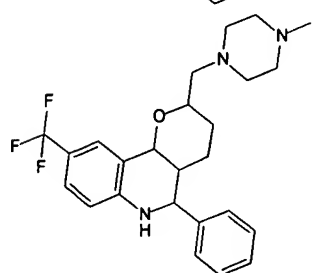
324 5.39 A

(145)



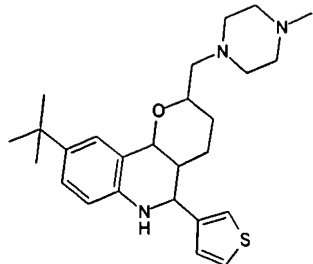
407 3.65 A

(146)



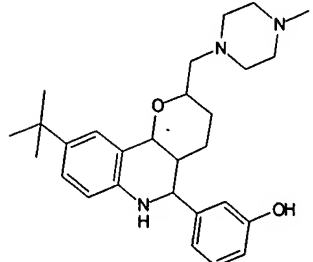
447 3.76 / 4.03 A

(147)



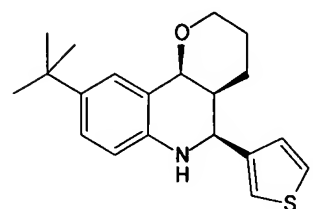
441 3.73 / 3.79 A

(148)



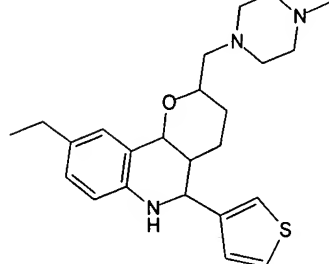
451 3.49 A

(149)



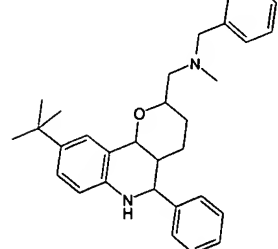
328 6.08 A

(150)



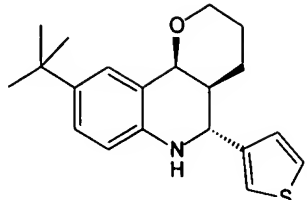
413 3.52 A

(151)



456 1.836 / 1.930 C

(152)

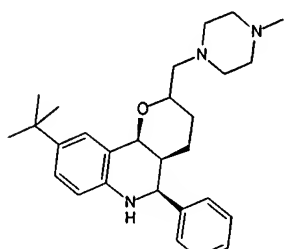


328

5.73

A

(153)

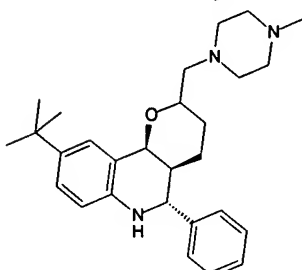


435

3.95

A

(154)

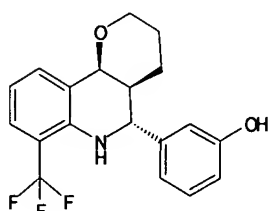


435

3.97

A

(155)

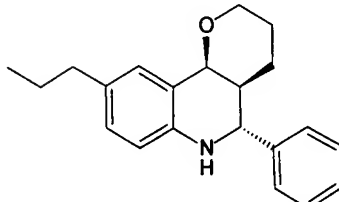


350

5.36

A

(156)

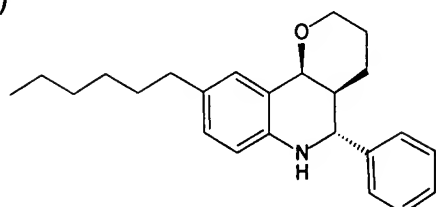


308

5.87

A

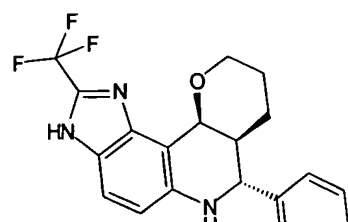
(157)



351

6.64

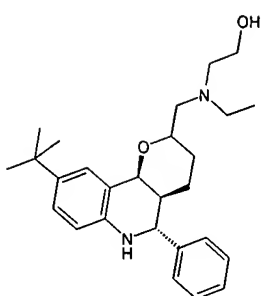
A



(158)

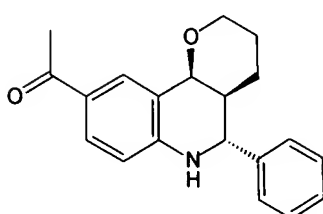
374 4.93 A

(159)



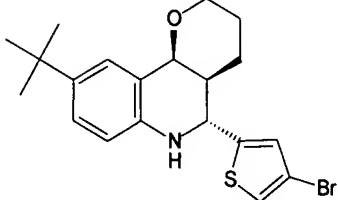
424 4.88 A

(160)



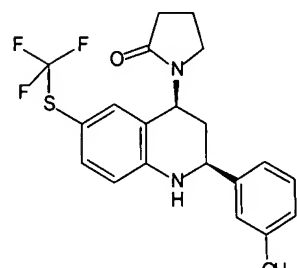
308 5.04 A

(161)



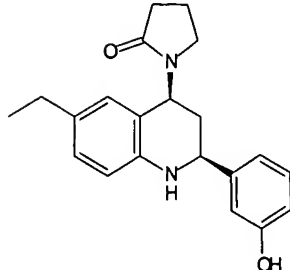
406/408 6.48 A

(162)



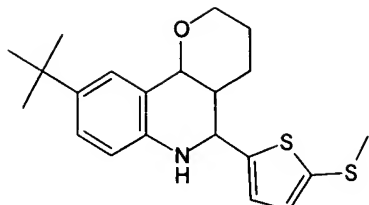
409 5.04 A

(163)



337 4.11 A

(164)

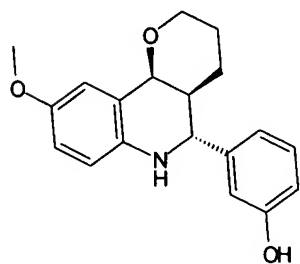


375

6.53

A

(165)

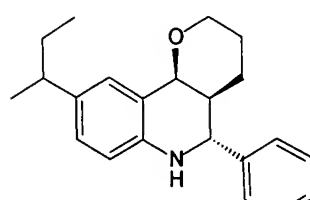


312

3.6

A

(166)

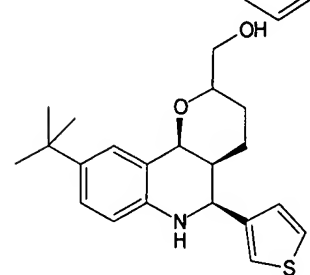


322

6.11

A

(167)

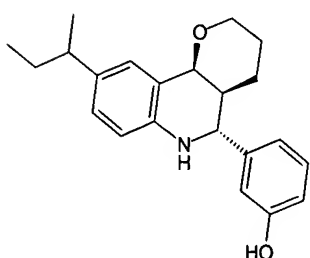


359

5.15

A

(168)

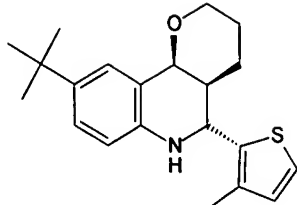


338

5.28

A

(169)

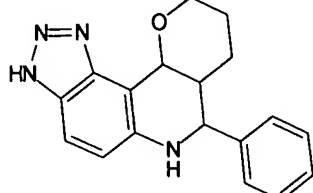


343

6.32

A

(170)

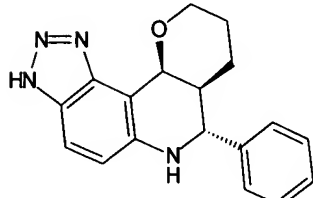


307

4.40 / 4.56

A

(171)

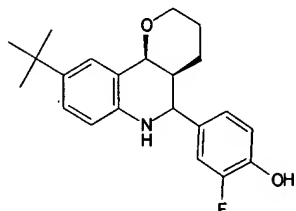


307

4.27

A

(172)

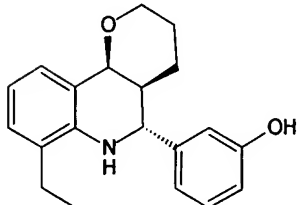


356

5.31

A

(173)



310

5.17

A

(174)

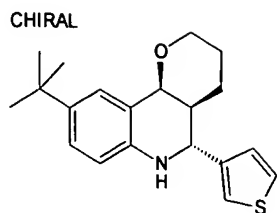


310

5.33

A

(175)

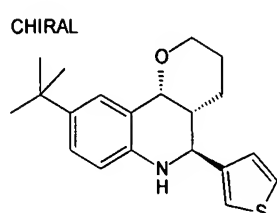


328

5.71

A

(176)

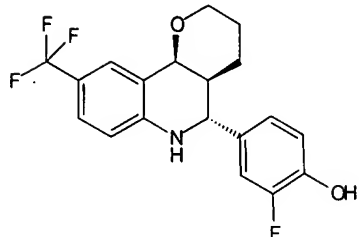


328

5.68

A

(177)

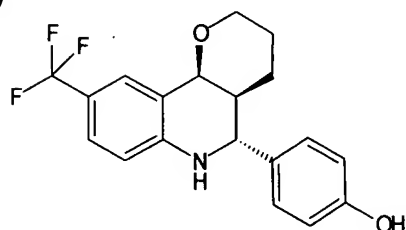


368

5.25

A

(178)

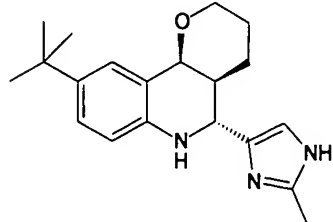


350

5.2

A

(179)

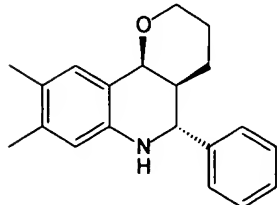


326

3.97

A

(180)

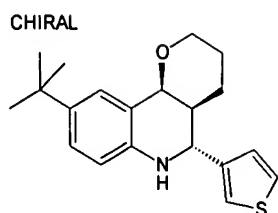


294

5.2

A

(181)

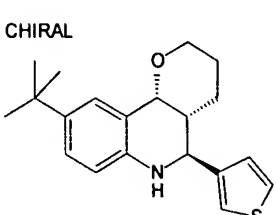


328

5.01

A

(182)

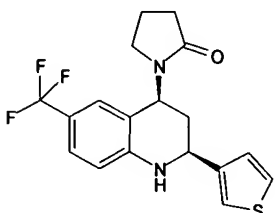


328

5.01

A

(183)

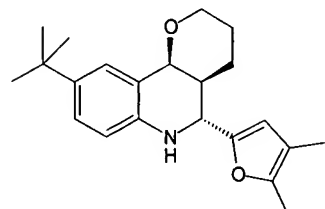


367

2.142

C

(184)

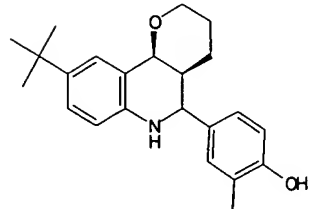


340

2.992

C

(185)

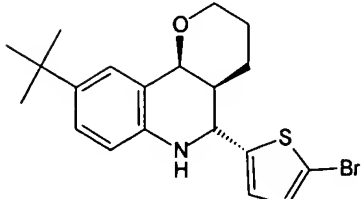


352

4.96

A

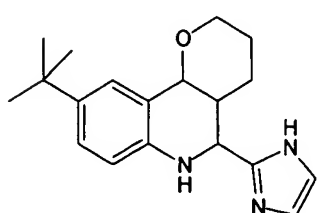
(186)



406/408

6.61

A



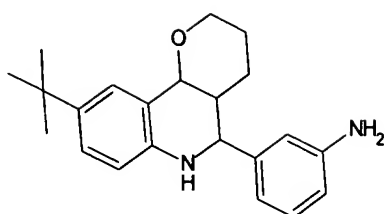
(187)

312

1.213

C

(188)

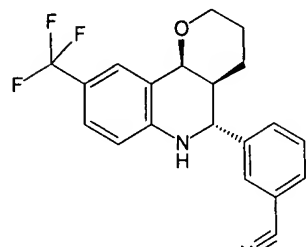


337

4.32

A

(189)

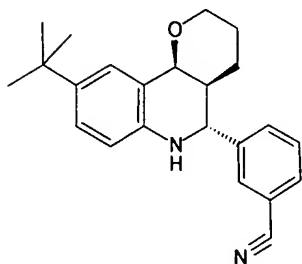


359

2.403

C

(190)

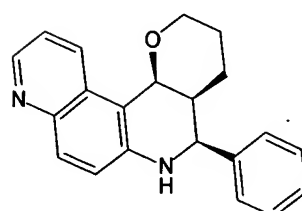


347

2.921

C

(191)

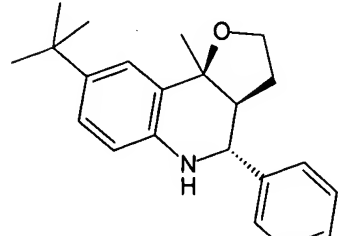


317

4.05

A

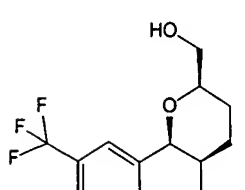
(192)



322

6.53 ?

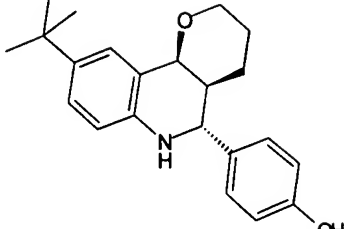
A



(193)

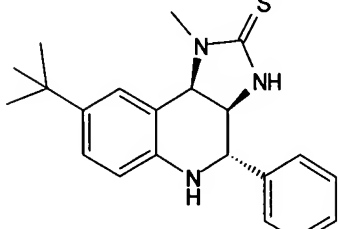
380 4.77 A

(194)



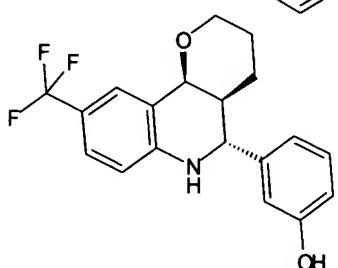
338 4.91 A

(195)



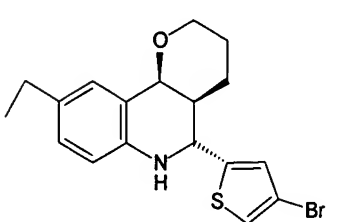
353 5.65 A

(196)



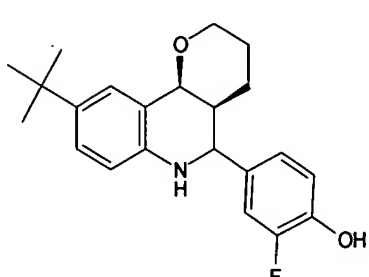
350 5.31 A

(197)

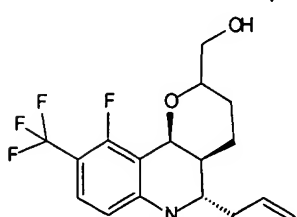


378/380 6.21 A

(198)



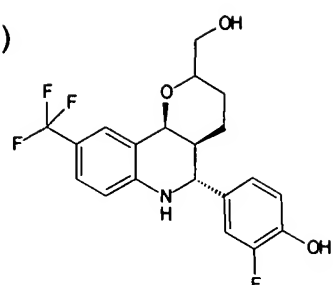
356 5.69 B



(199)

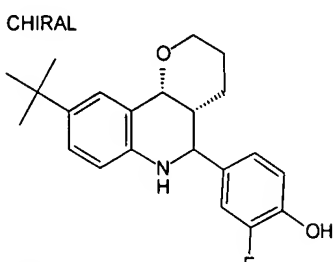
400 5.49 A

(200)



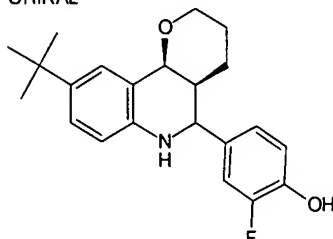
398 4.69/ 4.85 A

(201)



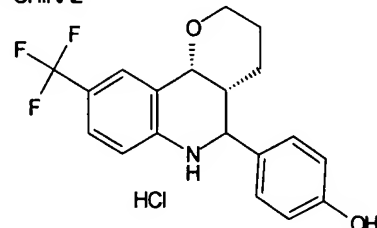
356 5.73 B

(202)



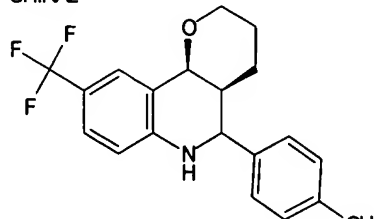
356 5.73 B

(203)

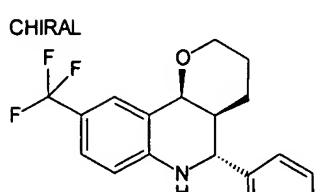


387 5.69 B

(204)



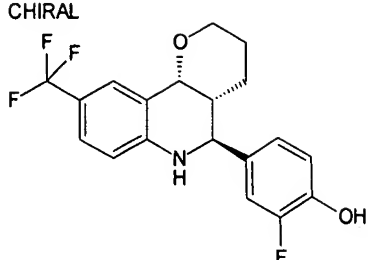
350 5.67 B



(205)

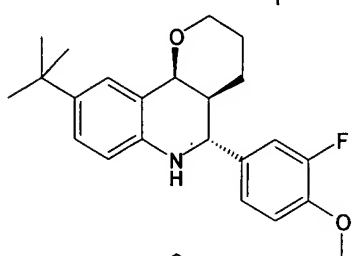
368 5.41 A

(206) CHIRAL



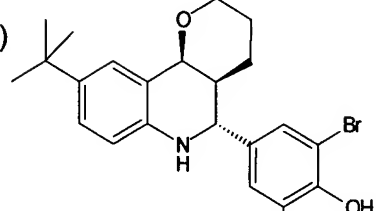
368 5.44 A

(207)

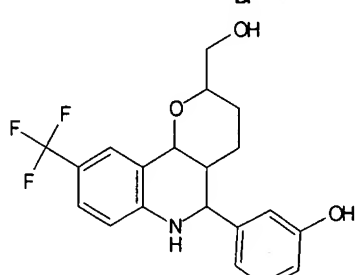


370 6.08 A

(208)

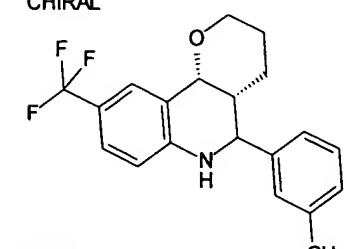
494/496/49
8 6.19 A

(209)

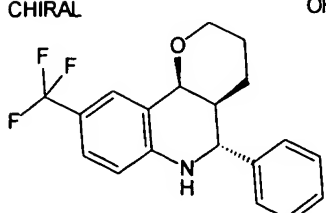


417 5.02 B

(210) HCl CHIRAL



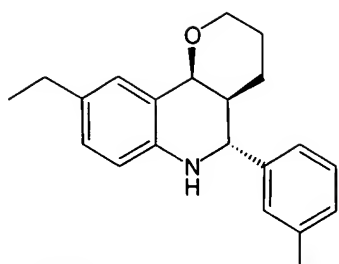
350 5.33 A



(211)

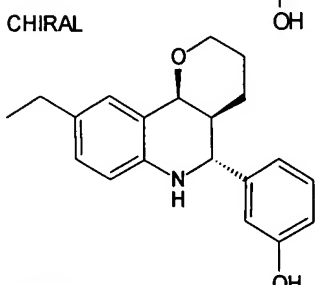
350 5.33 A

(212)



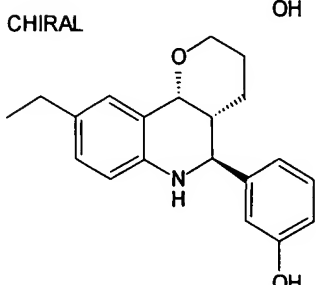
310 4.75 A

(213) CHIRAL



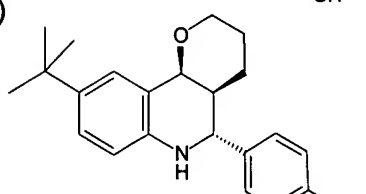
310 4.75 A

(214) CHIRAL



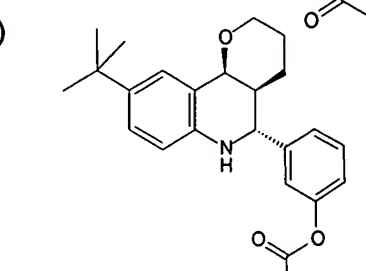
310 4.72 A

(215)

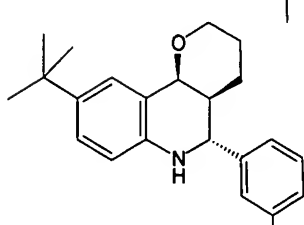


380 5.07 A

(216)



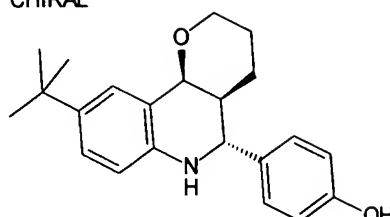
381 6 A



(217)

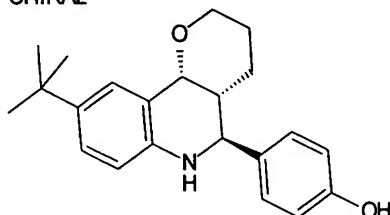
357 4.4 A

(218) CHIRAL



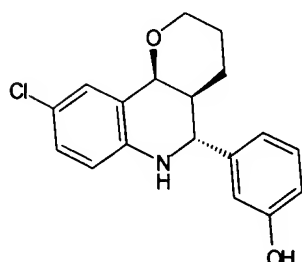
338 4.99 A

(219) CHIRAL



338 4.99 A

(220)



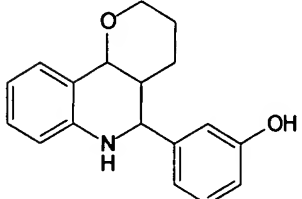
317 5.2 A

(221)



317 5.36 A

(222)



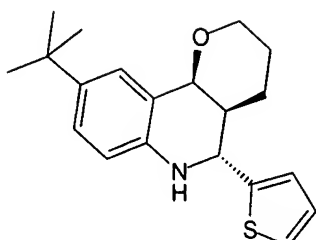
282 4.53/ 4.88 A



(223)

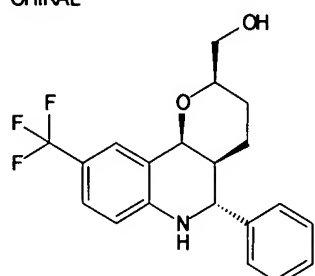
383 6.03/ 6.19 A

(224)



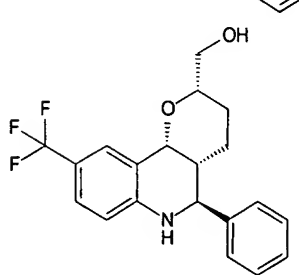
328 6.59 B

(225) CHIRAL



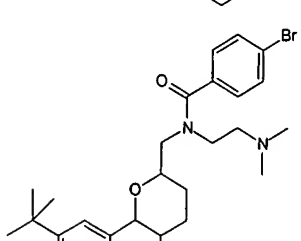
364 5.6 A

(226)



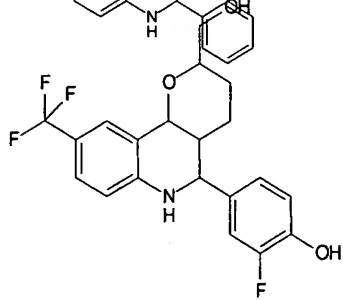
364 5.6 A

(227)



605/607 5.47 A

(228)



398 4.59 A

(229)

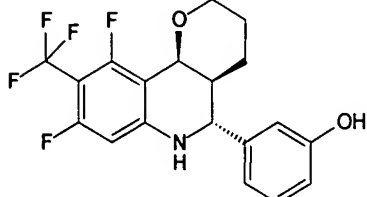


354

4.51

A

(230)

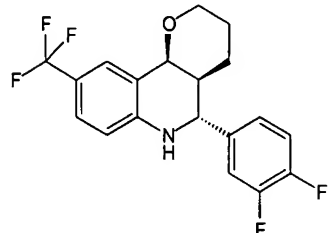


386

5.49

A

(231)

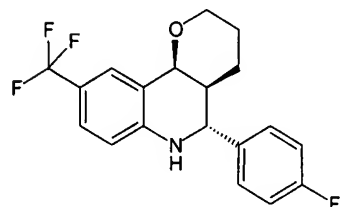


370

6.11

A

(232)

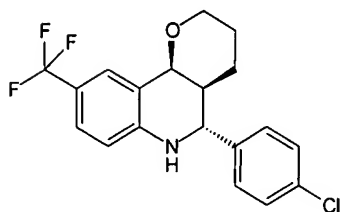


352

6.03

A

(233)

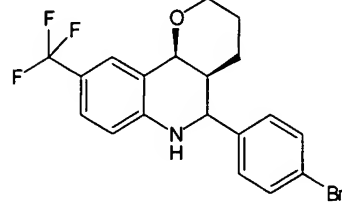


369

6.29

A

(234)

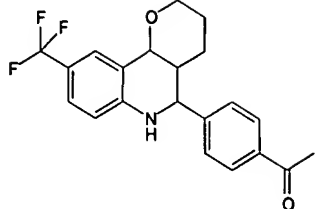


412/414

6.35

A

(235)

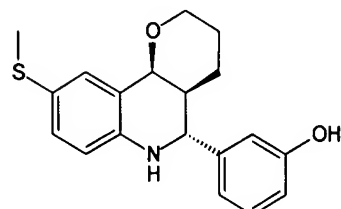


376

5.73 / 5.81

A

(236)

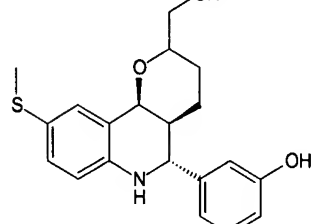


328

4.93

A

(237)

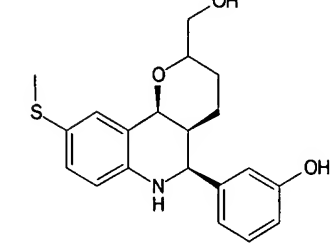


358

4.21

A

(238)

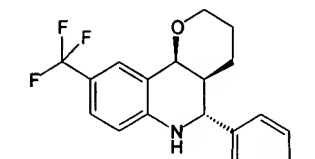


358

4.16

A

(239)

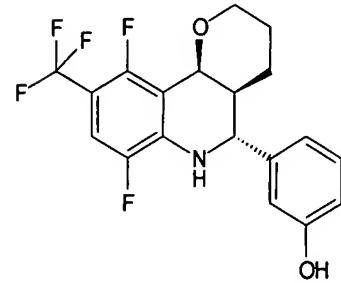


348

6.27

A

(240)

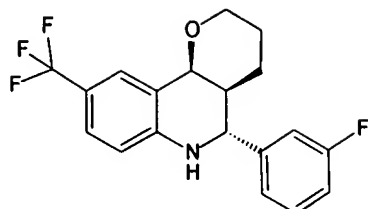


386

5.52

A

(241)

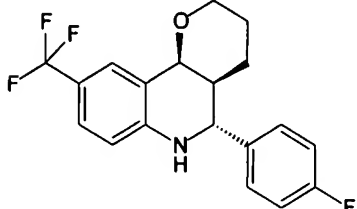


352

6.05

A

(242)

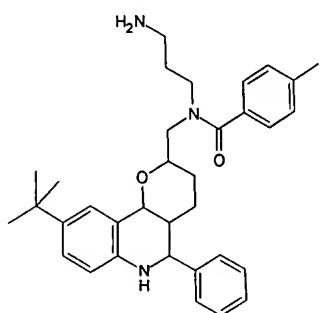


352

6.08

A

(243)

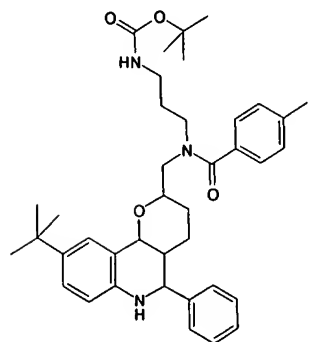


527

1.777 / 1.902

C

(244)

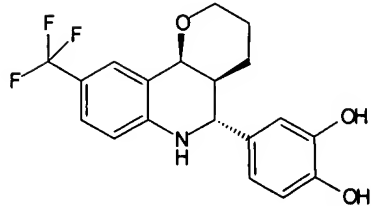


627

2.770 / 2871

C

(245)

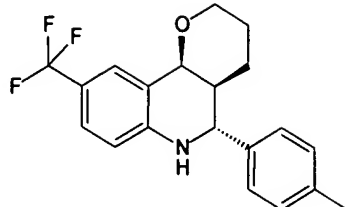


366

4.99

A

(246)

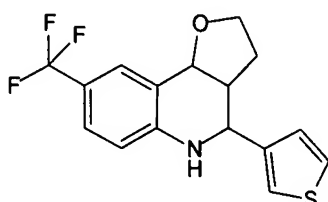


348

6.29

A

(247)

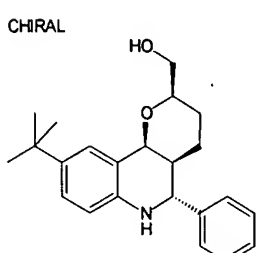


326

2.242

C

(248)

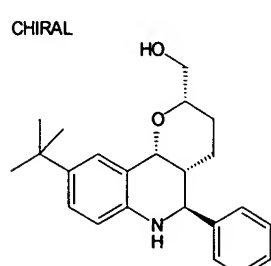


352

5.44

A

(249)

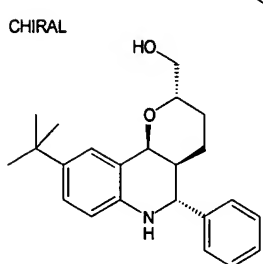


352

5.44

A

(250)

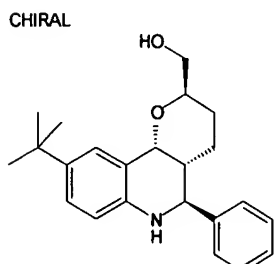


352

5.7

A

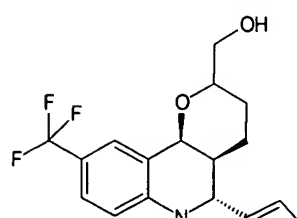
(251)



352

5.7

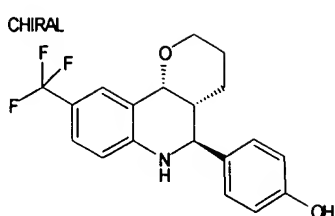
A



(252)

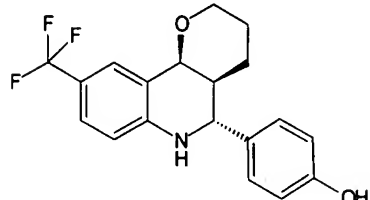
370 5.36 A

(253)



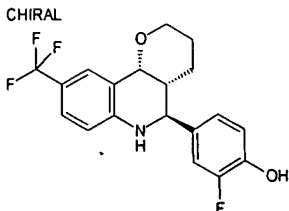
350 5.65 B

(254)



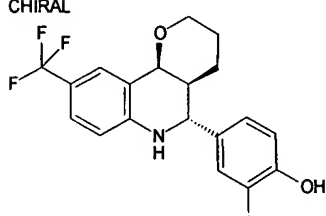
350 5.67 B

(255)



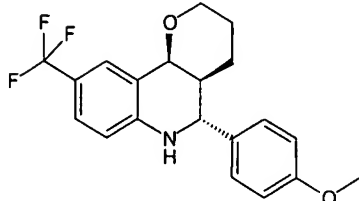
368 5.52 A

(256)

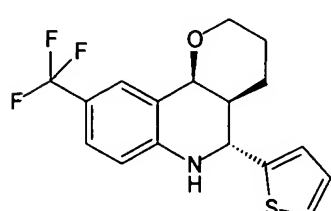


368 5.52 A

(257)



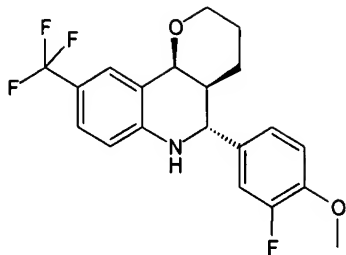
364 6.03 A



(258)

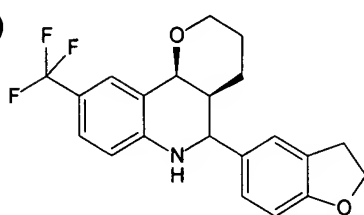
340 5.97 A

(259)



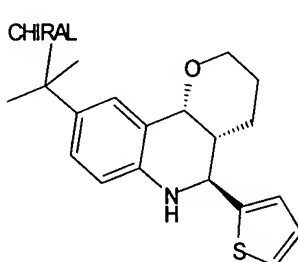
382 6 A

(260)



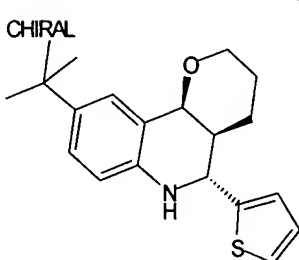
376 6 A

(261)



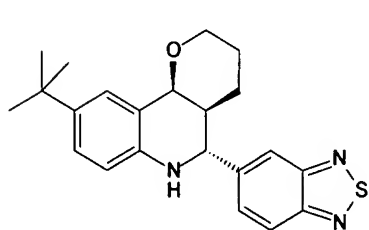
328 6.5 B

(262)



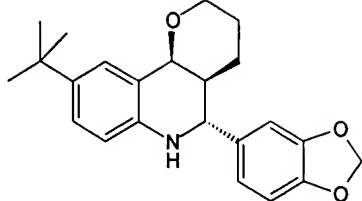
328 6.53 B

(263)



381 6.29 A

(264)

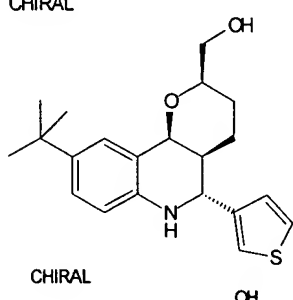


366

5.87

A

(265) CHIRAL

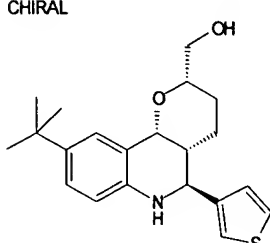


359

5.15

B

(266) CHIRAL

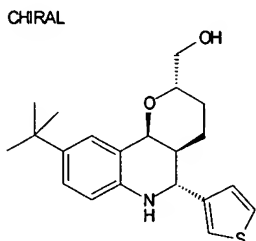


359

5.15

B

(267) CHIRAL

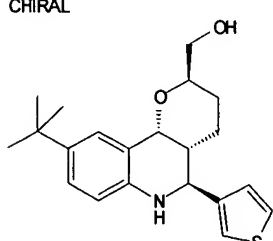


359

5.46

B

(268) CHIRAL

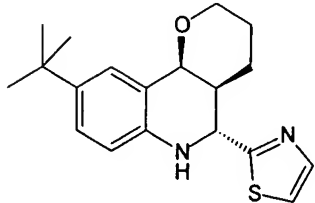


359

5.47

B

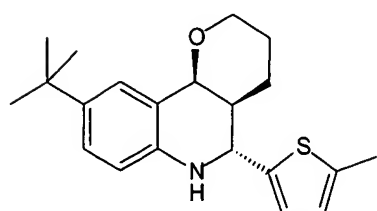
(269)



329

5.57

A



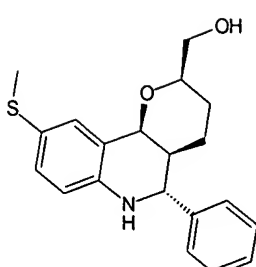
(270)

343

6.4

A

(271)

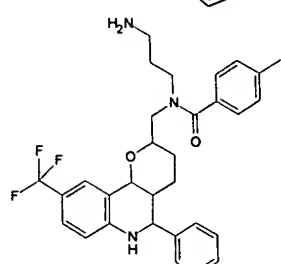


342

5.04

A

(272)

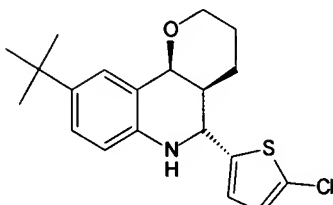


539

4.88 / 5.09

A

(273)

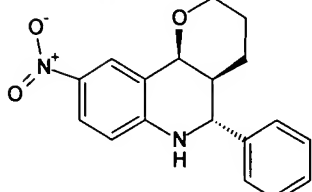


363

6.69

A

(274)

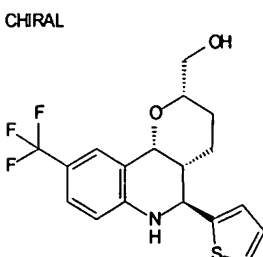


311

5.47

A

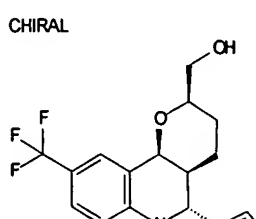
(275)



370

5.36

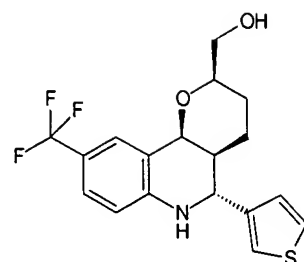
A



(276)

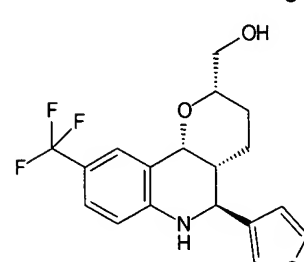
370 5.36 A

(277)



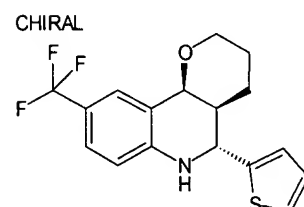
370 5.52 A

(278)



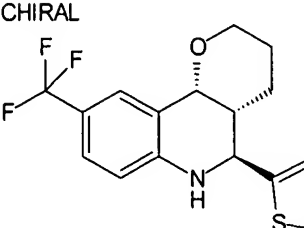
370 5.52 A

(279)



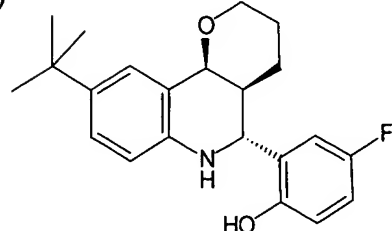
340 5.99 A

(280)



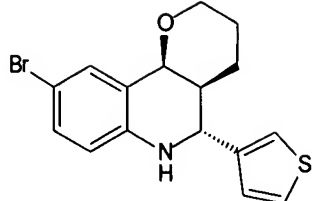
340 6.00 A

(281)



356 5.57 A

(282)

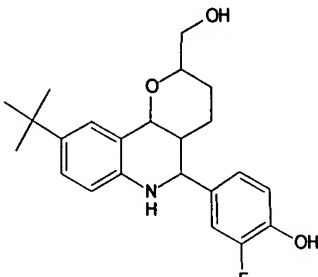


350/352

5.87

A

(283)

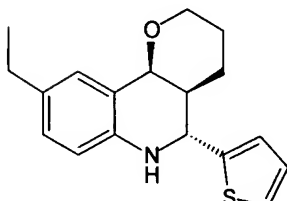


386

4.59 / 4.72

A

(284)

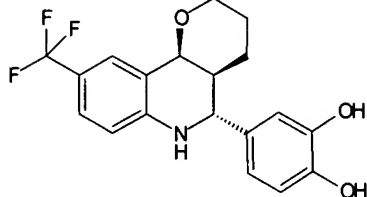


300

5.79

A

(285)

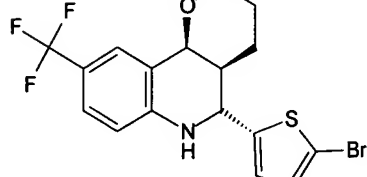


366

4.93

A

(286)

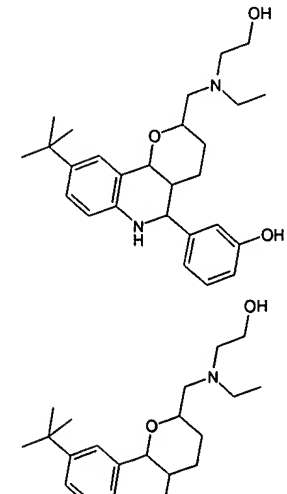


418/420

6.35

A

(287)



440

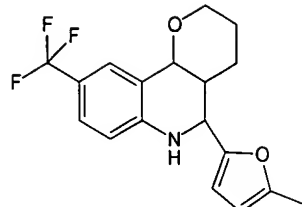
1.378

A

(288)

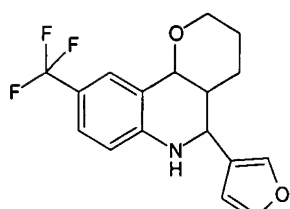
458 4.27 A

(289)



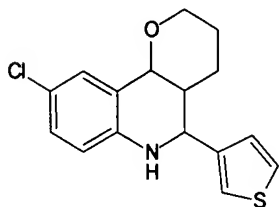
338 2.409 C

(290)



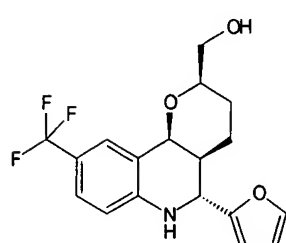
324 5.65 A

(291)



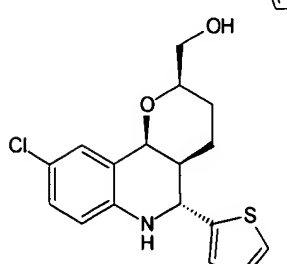
307 2.32 C

(292)

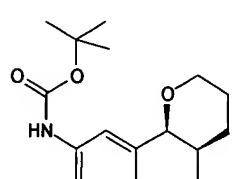


354 1.903 C

(293)



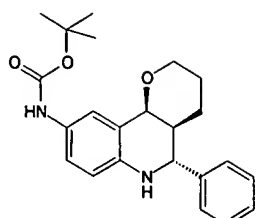
337 5.12 A



(294)

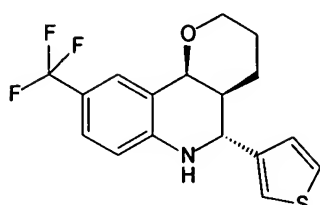
381 5.55 A

(295)



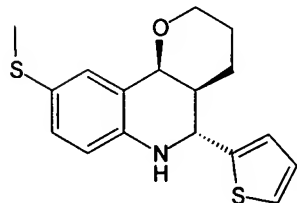
381 5.2 A

(296)



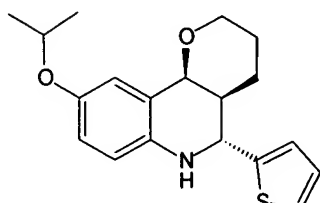
340 2.401 C

(297)



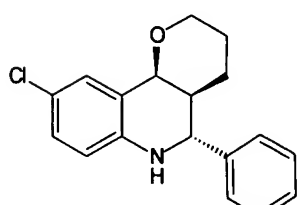
318 2.259 C

(298)

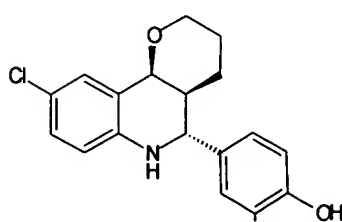


330 2.07 C

(299)



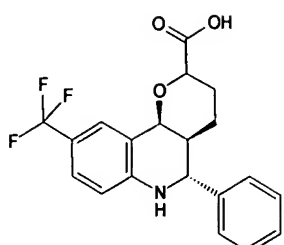
301 2.416 C



(300)

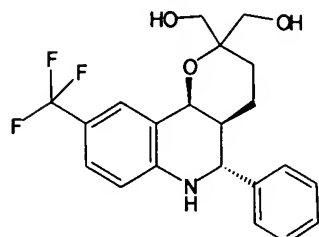
335 2.013 C

(301)



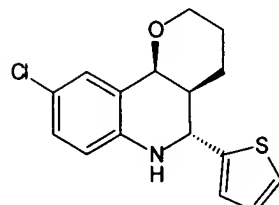
378 1.922 / 2.081 C

(302)



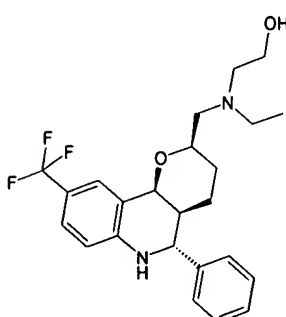
394 4.99 A

(303)



307 2.367 C

(304)



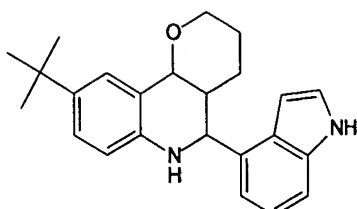
436 4.95 B

(305)		436	4.78	B
(306)		300	5.8	A
(307)		300	5.8	A
(308)		328	4.88	A
(309)		545	6.32	A

(310)

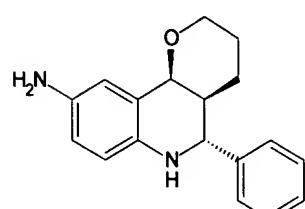
368 4.66 B

(311)



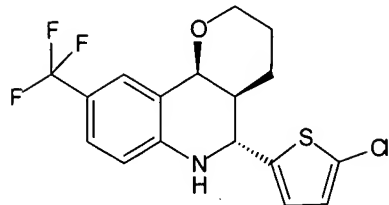
362 5.57 A

(312)



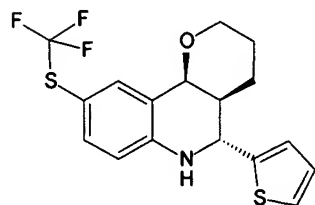
281 3.63 A

(313)



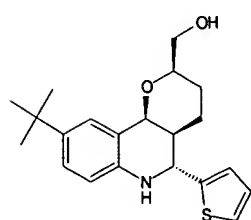
375 6.46 A

(314)



372 6.32 A

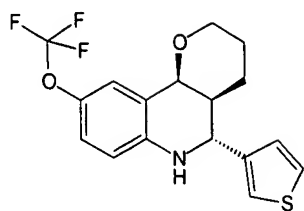
(315)



359 5.52 A

(316)		372	2.602	C
(317)		310	4.93 / 5.07	A
(318)		328	6.53	A
(319)		359	5.5	A
(320)		359	5.49	A
(321)		324	5.84	A

(322)

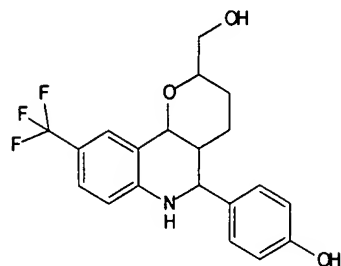


356

6.11

A

(323)

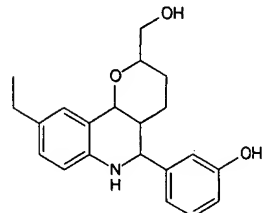


380

4.56 / 4.72

A

(324)

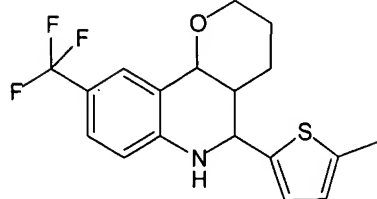


340

3.84 / 4.10

A

(325)

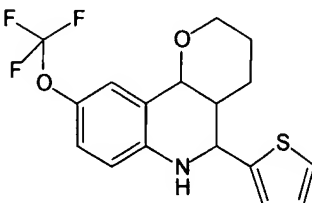


354

6.32

A

(326)



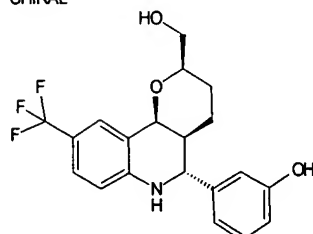
356

6.13

A

(327)

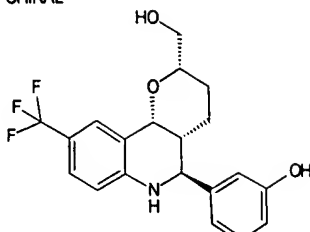
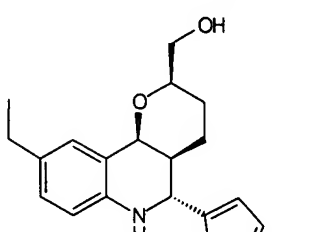
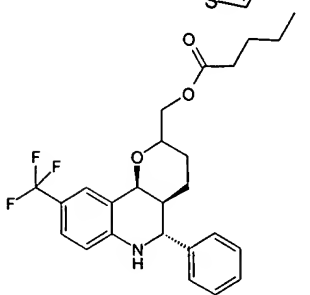
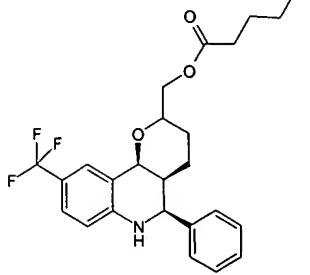
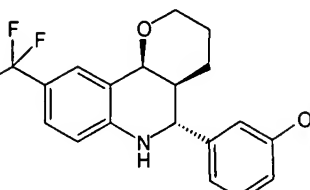
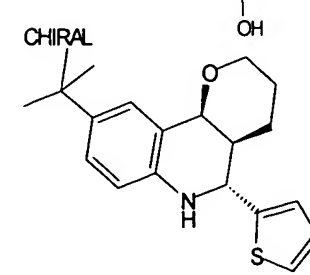
CHIRAL

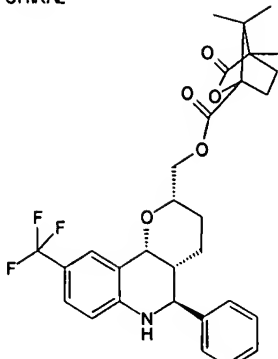
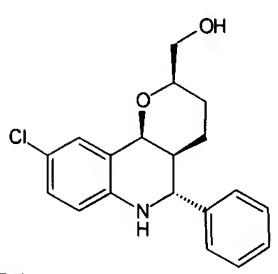
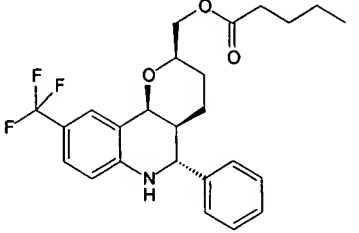
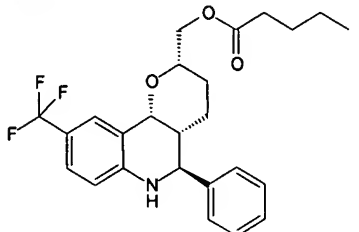
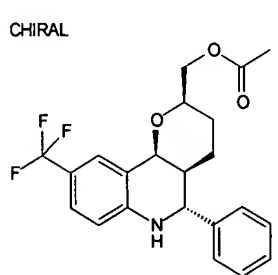


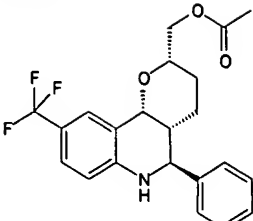
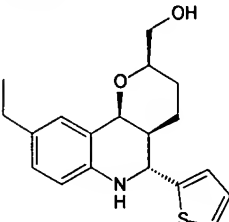
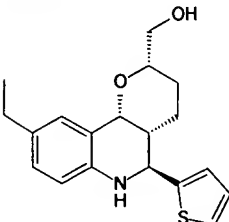
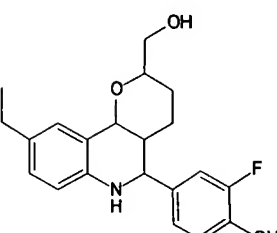
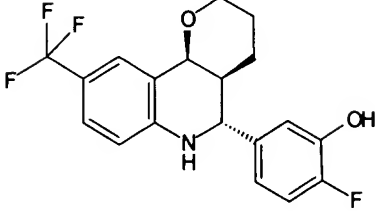
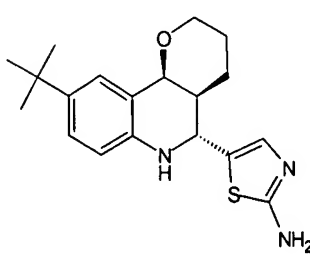
380

4.72

A

(328)	CHIRAL		380	4.72	A
(329)			330	5.12	A
(330)			449	6.69	A
(331)			449	6.59	A
(332)			366	4.83	A
(333)	CHIRAL		328	6.53	A

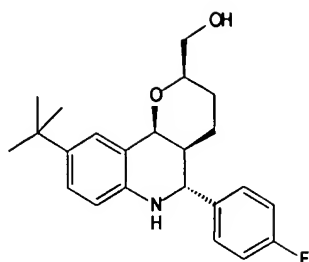
(334)	CHIRAL			
		545	6.32	A
(335)				
		331	5.25	A
(336)	CHIRAL			
		449	6.64	A
(337)	CHIRAL			
		449	6.64	A
(338)	CHIRAL			
		406	6.03	A

(339)	CHIRAL		406	6.03	A
(340)	CHIRAL		330	5.14	A
(341)	CHIRAL		330	5.13	A
(342)			358	4.16 / 4.37	A
(343)			368	1.998	C
(344)			344	4.43	A

- 139 -

(345)		406	6.19	A
(346)	CHIRAL 	375	6.46	A
(347)	CHIRAL 	375	6.46	A
(348)		441	6.21	A
(349)		378	6.08	A
(350)		382	5.49	A

(351)

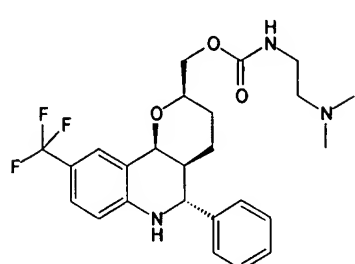


370

5.49

A

(352)

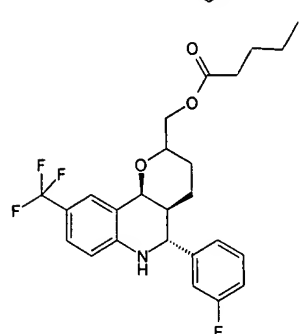


479

4.77

A

(353)

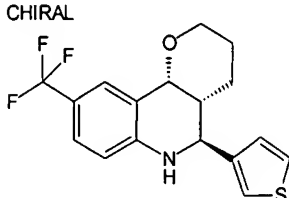


466

6.93

B

(354)

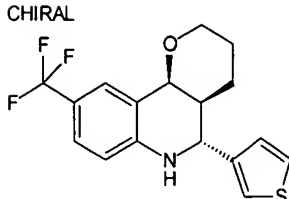


340

5.93

A

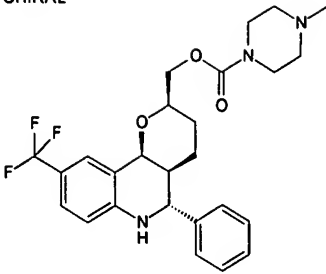
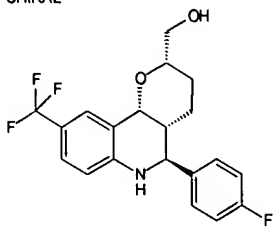
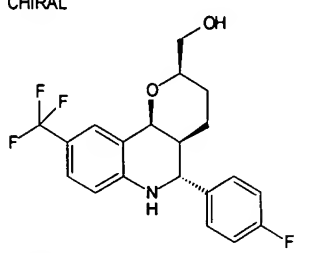
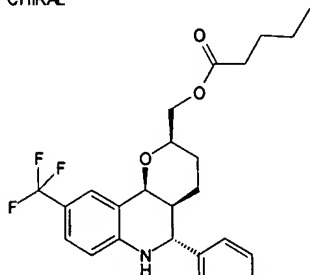
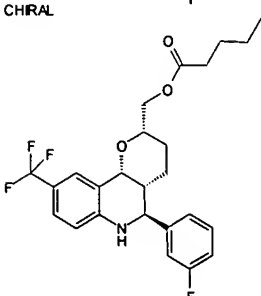
(355)

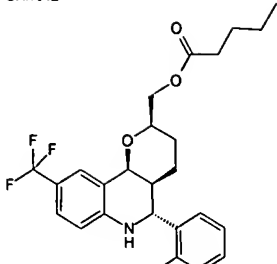
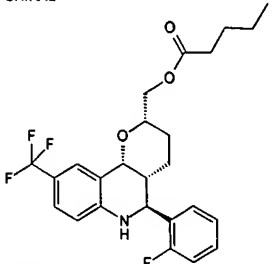
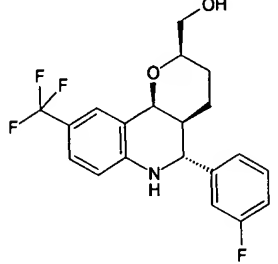
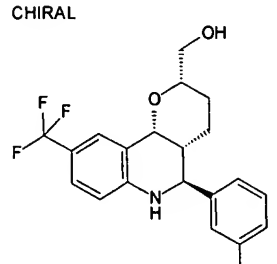
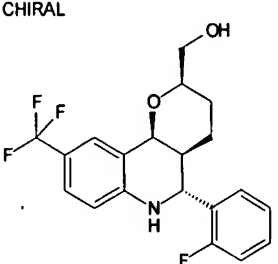


340

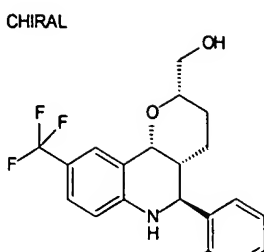
5.92

A

(356)	CHIRAL		491	4.83	A
(357)	CHIRAL		382	5.5	A
(358)	CHIRAL		382	5.51	A
(359)	CHIRAL		466	6.92	B
(360)	CHIRAL		466	6.93	B

(361)	CHIRAL		466	6.72	A
(362)	CHIRAL		466	6.72	A
(363)	CHIRAL		382	5.64	B
(364)	CHIRAL		382	5.64	B
(365)	CHIRAL		382	5.6	A

(366)

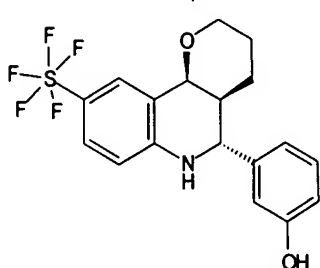


382

5.6

A

(367)

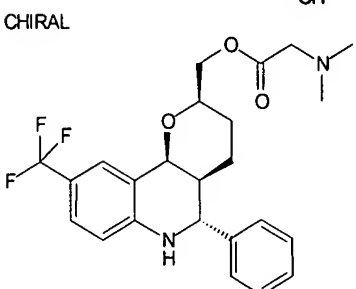


408

5.68

A

(368)

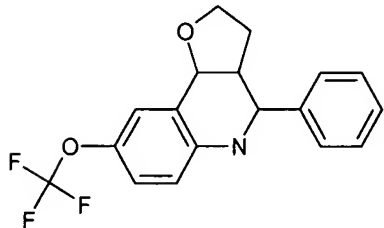


449

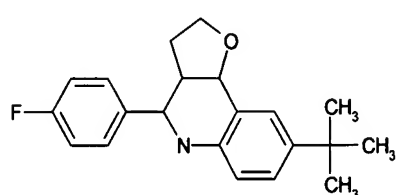
5.01

A

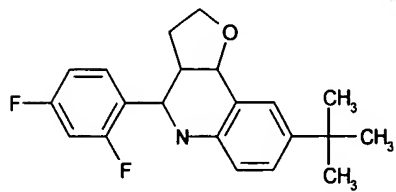
(369)



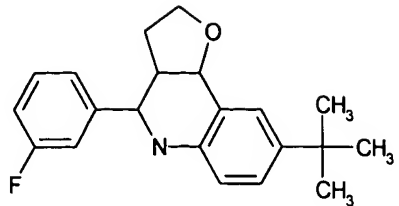
(370)



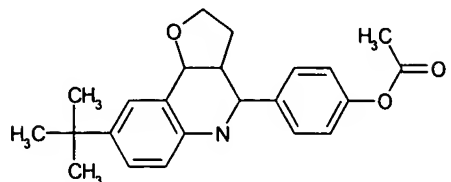
(371)



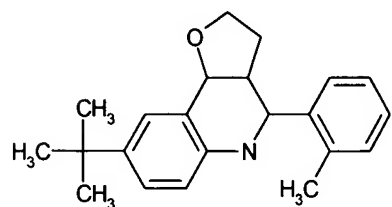
(372)



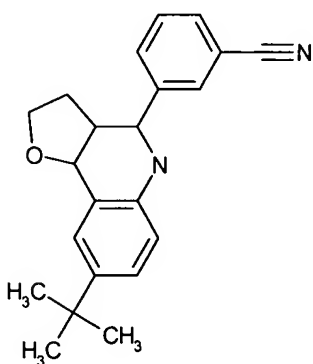
(373)



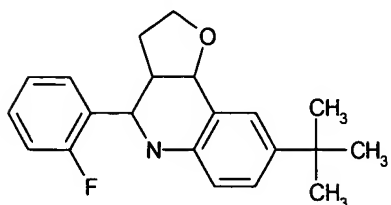
(374)



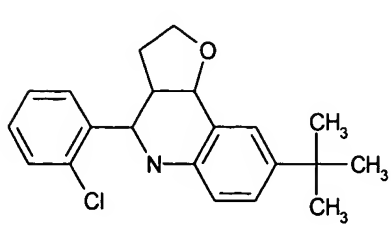
(375)



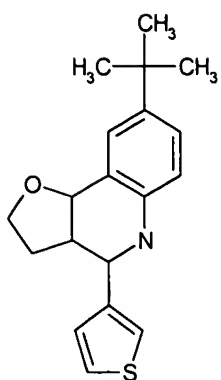
(376)



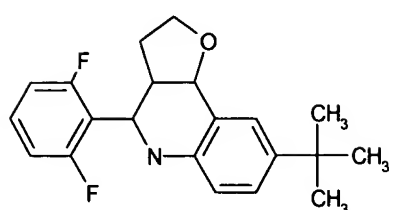
(377)



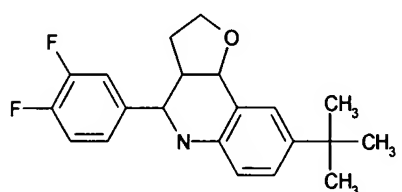
(378)



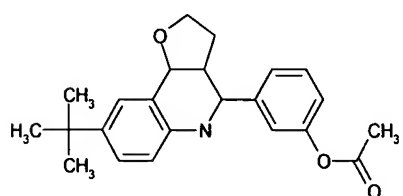
(379)



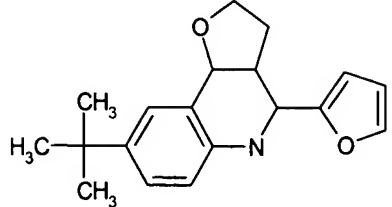
(380)



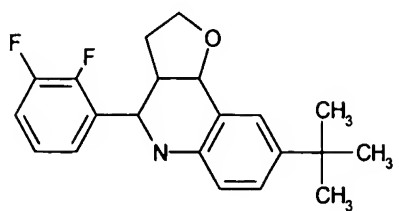
(381)



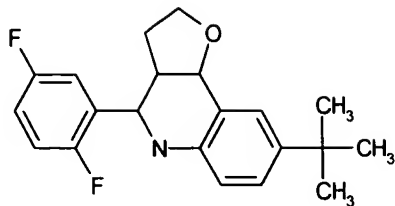
(382)



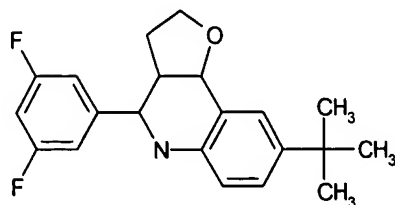
(383)



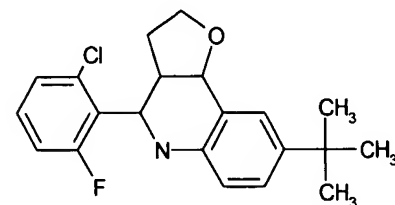
(384)



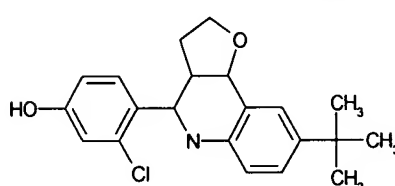
(385)



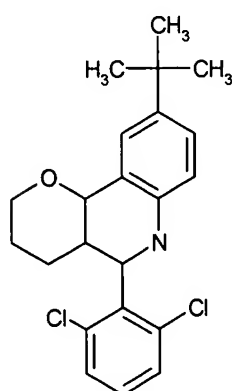
(386)



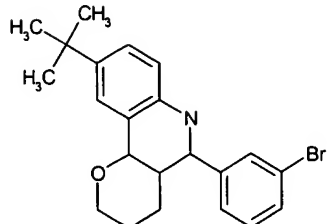
(387)

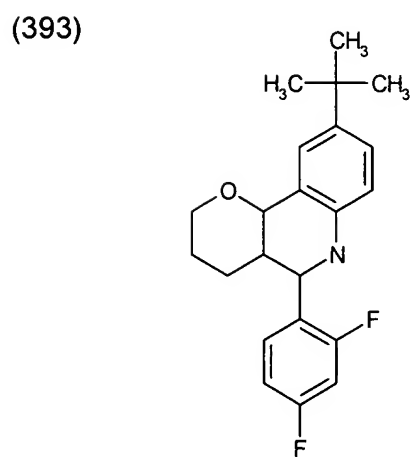
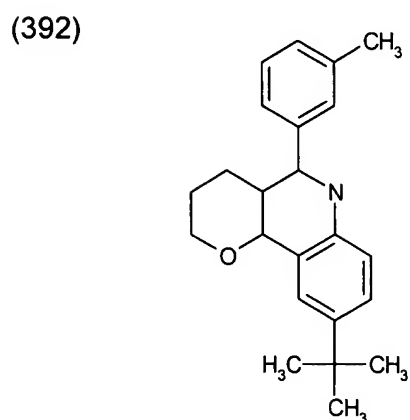
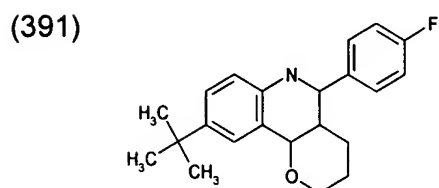
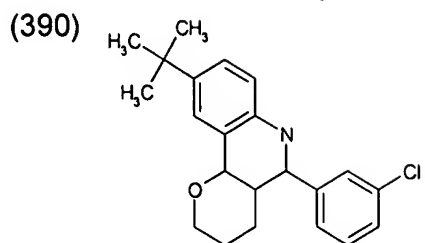


(388)

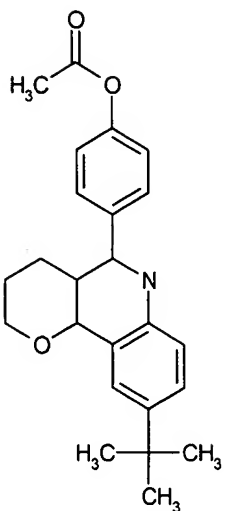


(389)

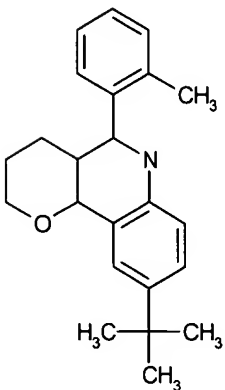




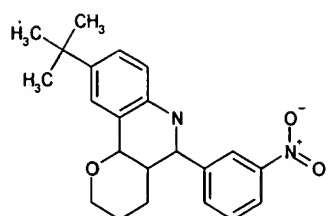
(394)



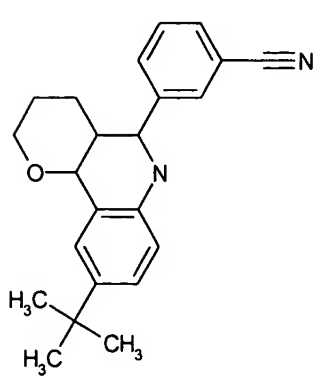
(395)



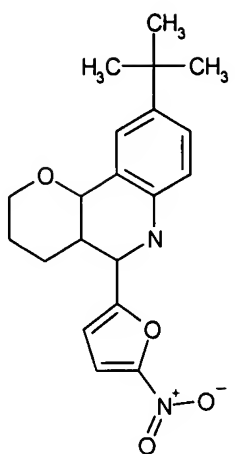
(396)



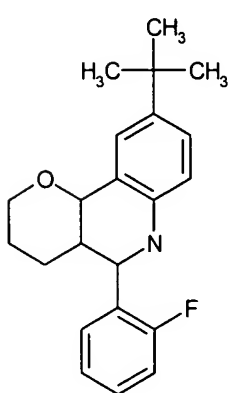
(397)



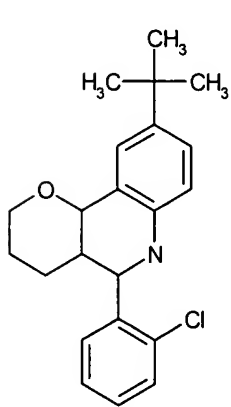
(398)



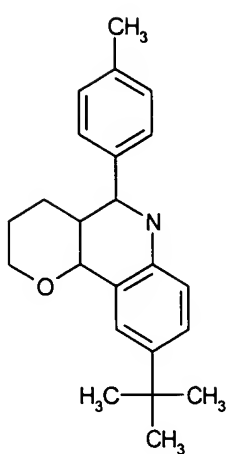
(399)



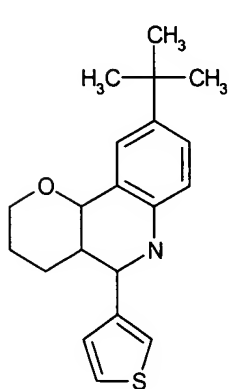
(400)



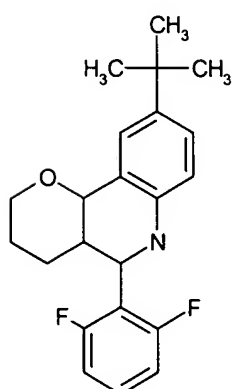
(401)



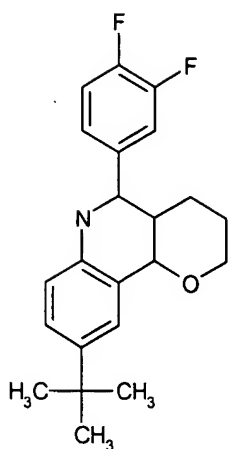
(402)



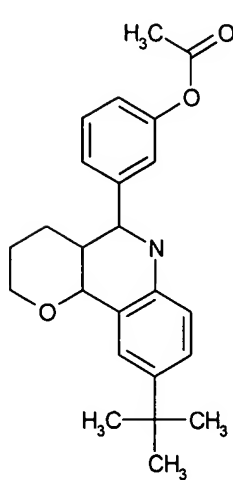
(403)



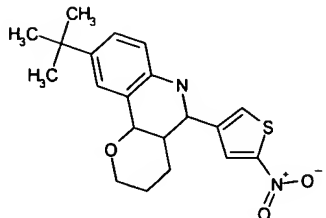
(404)



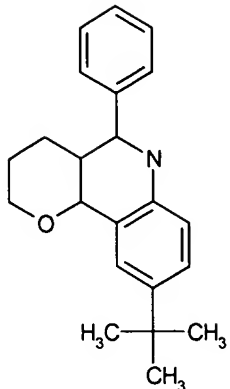
(405)



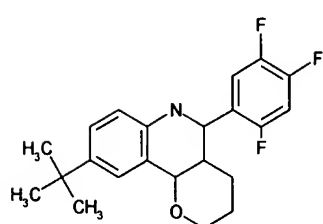
(406)



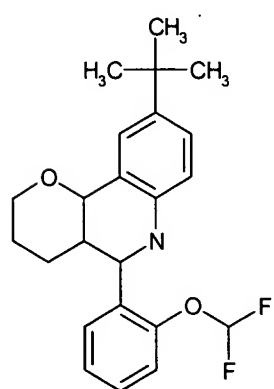
(407)



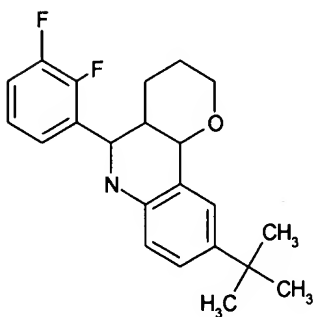
(408)



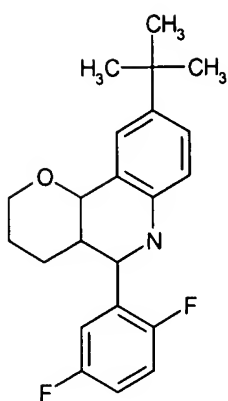
(409)



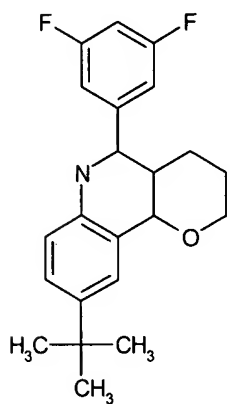
(410)



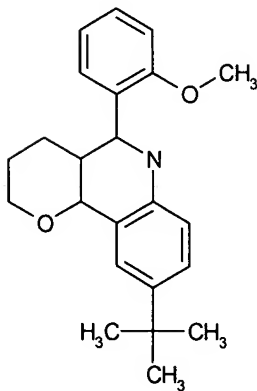
(411)



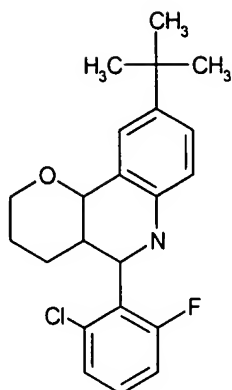
(412)



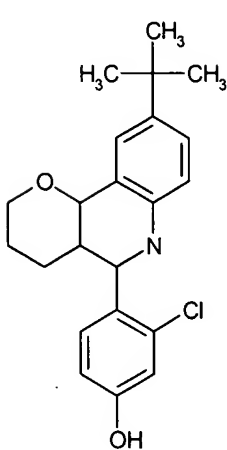
(413)



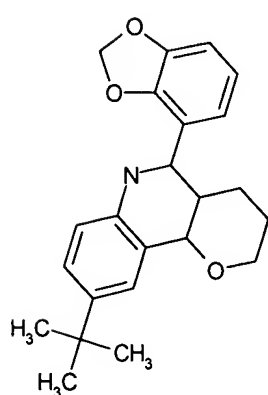
(414)



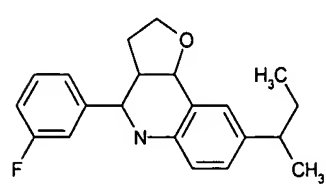
(415)



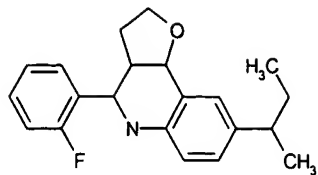
(416)



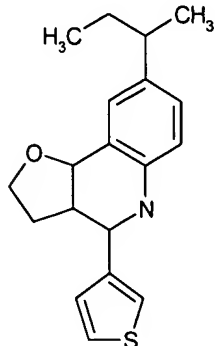
(417)



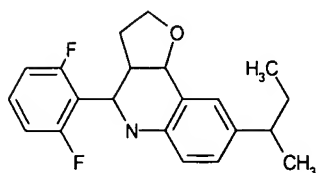
(418)



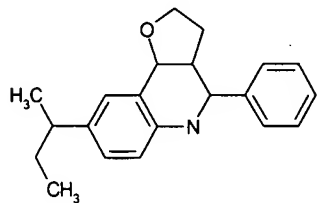
(419)



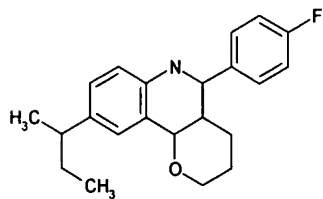
(420)



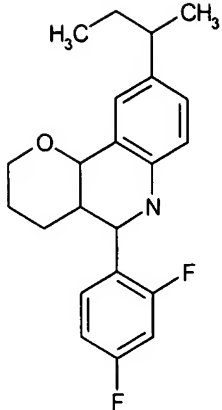
(421)



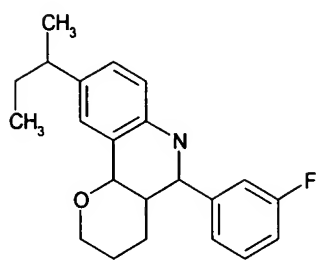
(422)



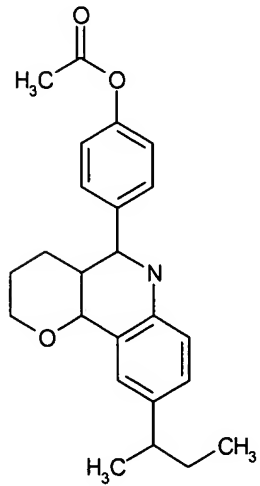
(423)



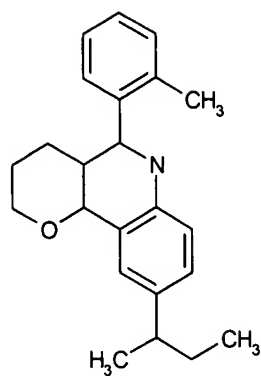
(424)



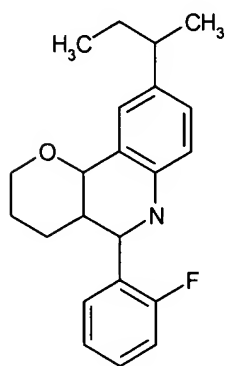
(425)



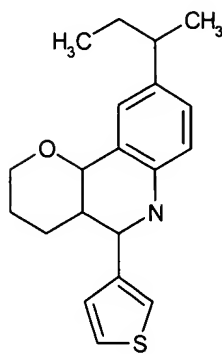
(426)



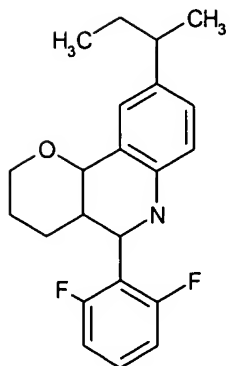
(427)



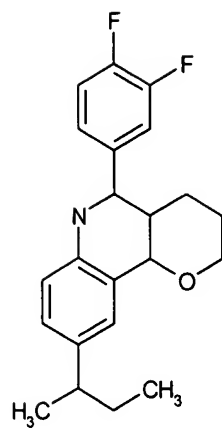
(428)



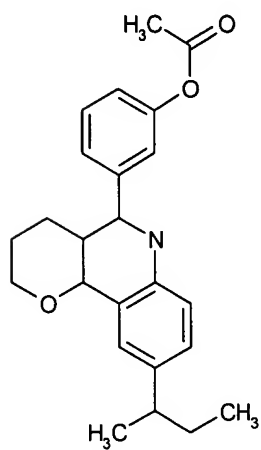
(429)



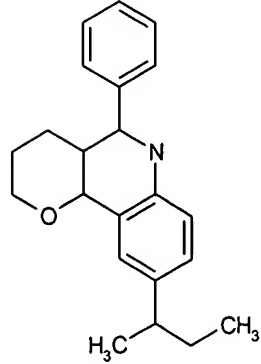
(430)



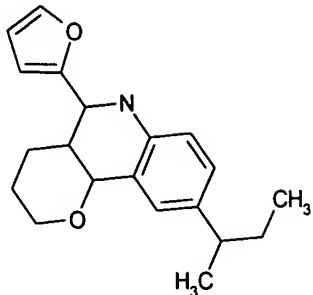
(431)



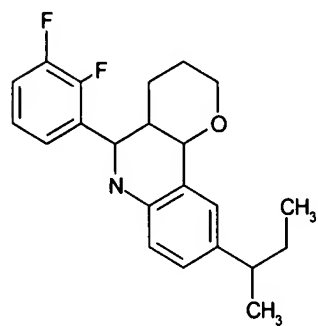
(432)



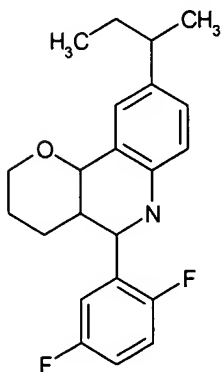
(433)



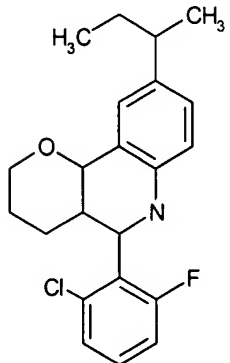
(434)



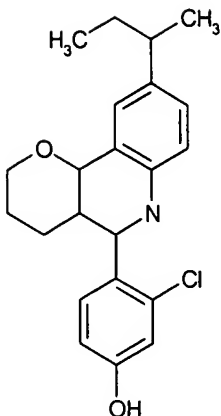
(435)



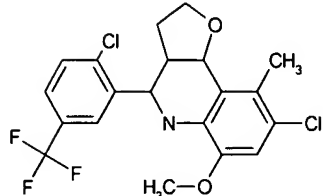
(436)



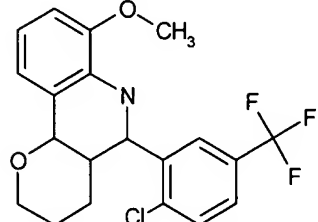
(437)



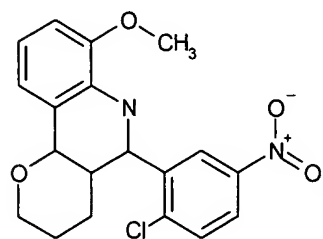
(438)



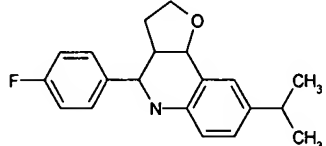
(439)



(440)



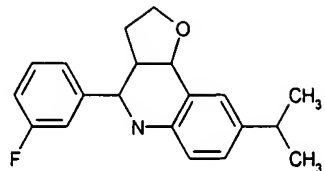
(441)



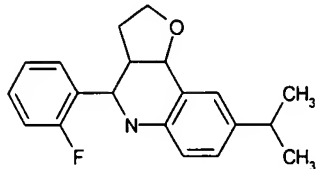
(442)



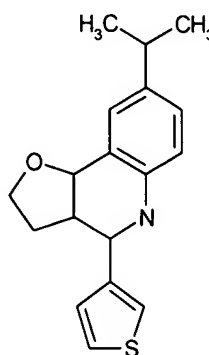
(443)



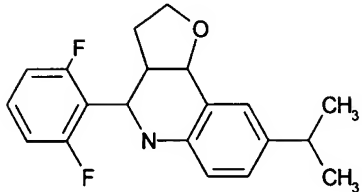
(444)



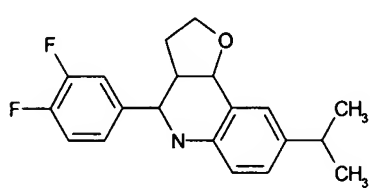
(445)



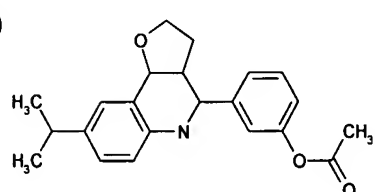
(446)



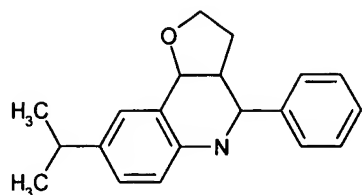
(447)



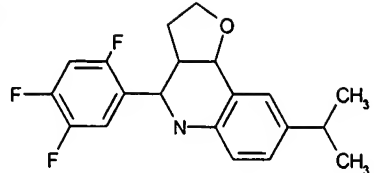
(448)



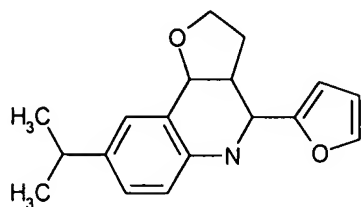
(449)



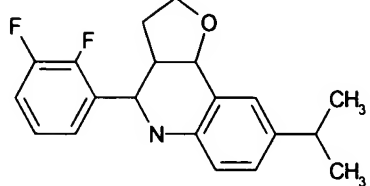
(450)



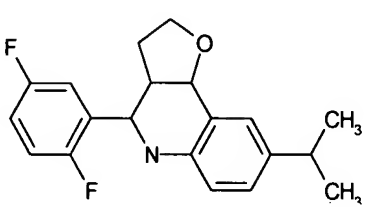
(451)



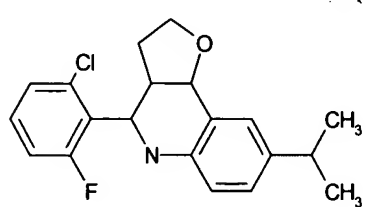
(452)



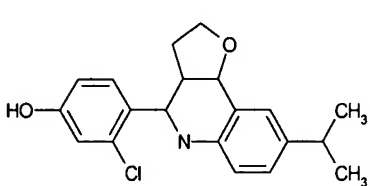
(453)



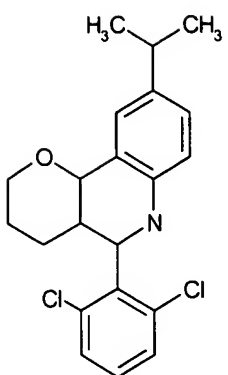
(454)



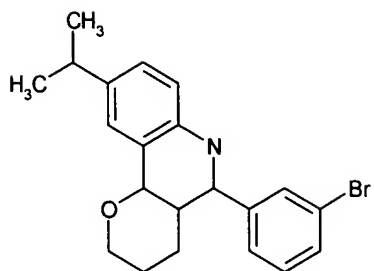
(455)



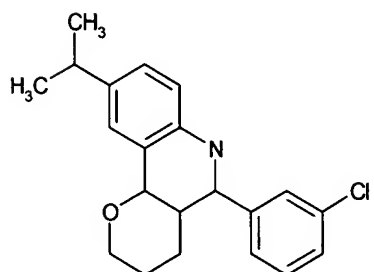
(456)



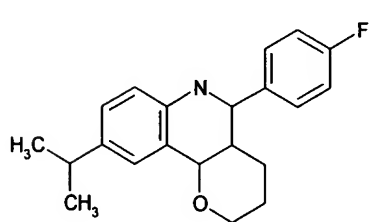
(457)



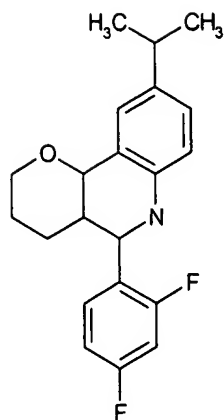
(458)



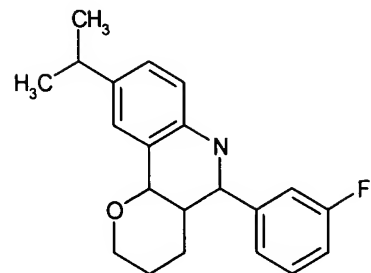
(459)



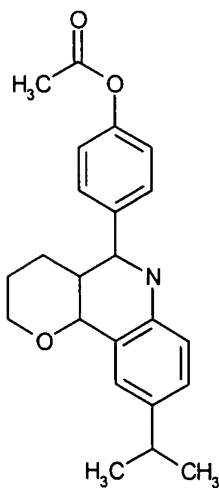
(460)



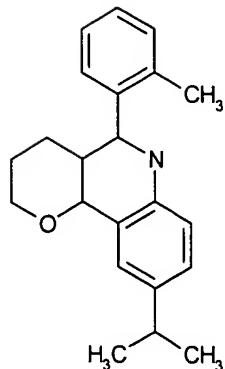
(461)

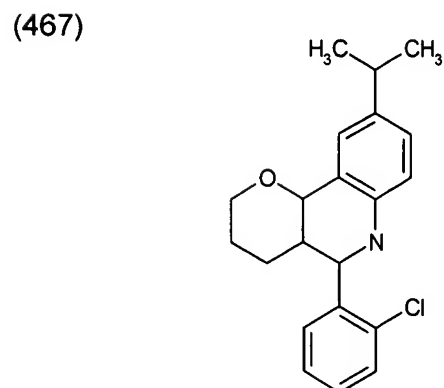
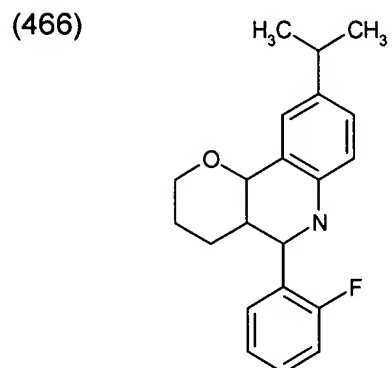
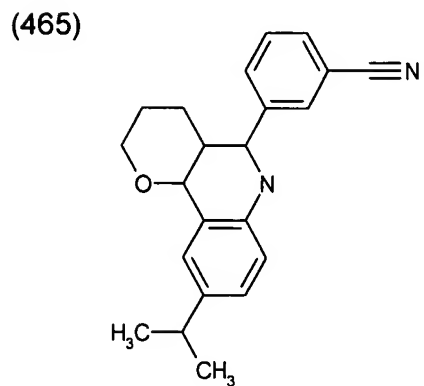
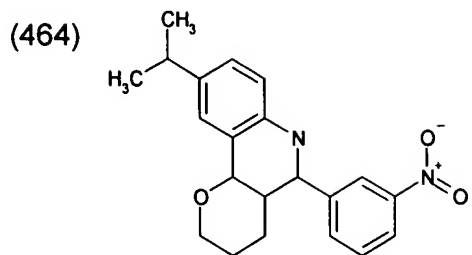


(462)

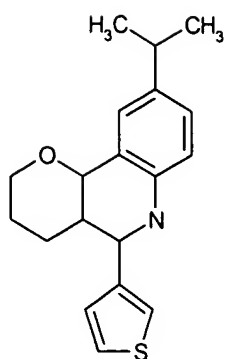


(463)

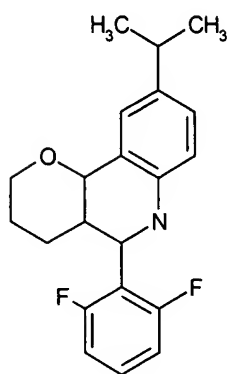




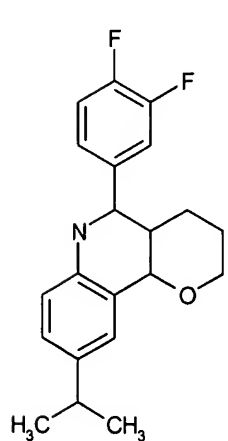
(468)



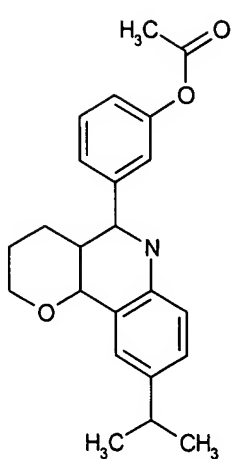
(469)



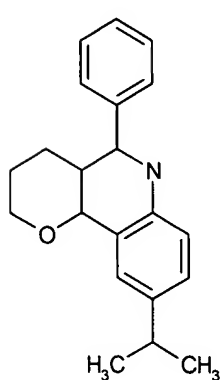
(470)



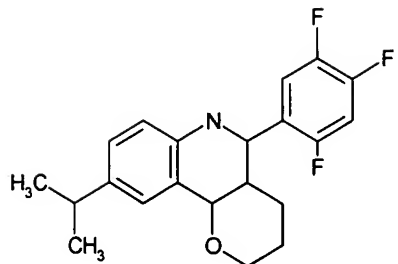
(471)



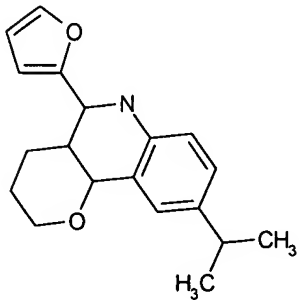
(472)



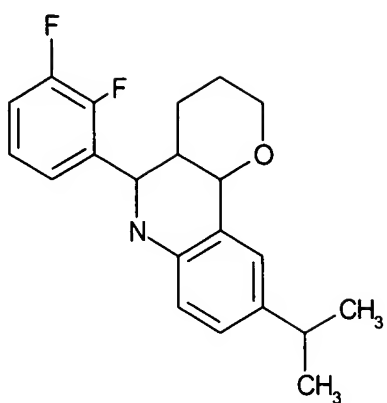
(473)



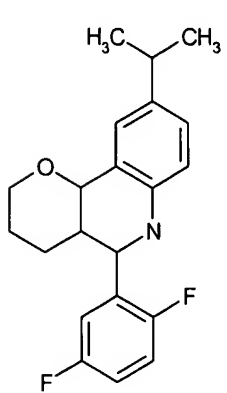
(474)



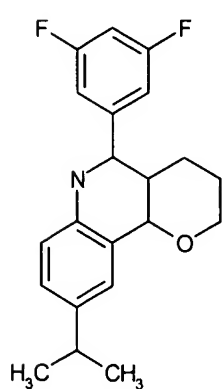
(475)



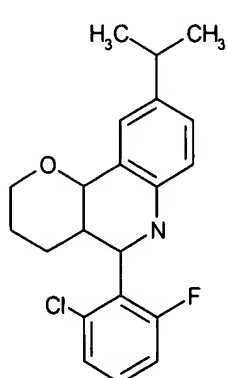
(476)



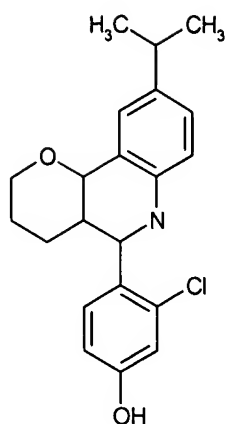
(477)



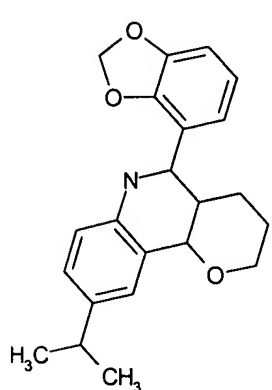
(478)



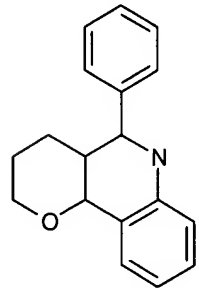
(479)



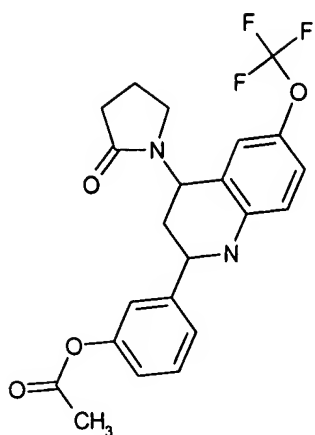
(480)



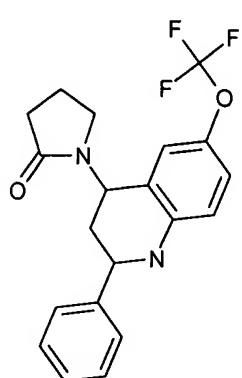
(481)



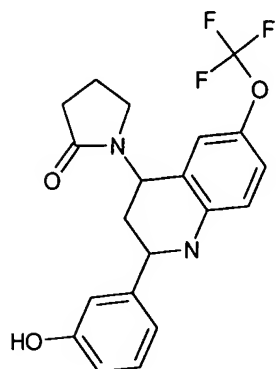
(482)



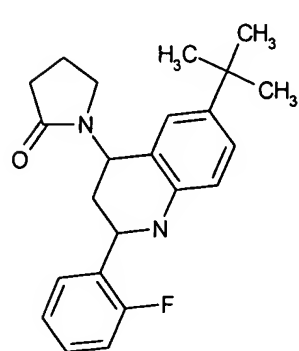
(483)



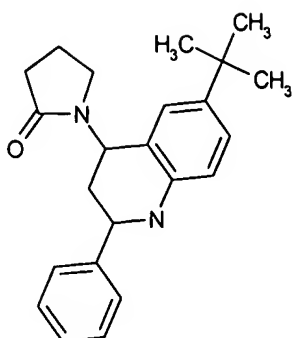
(484)



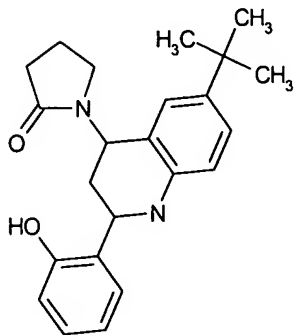
(485)



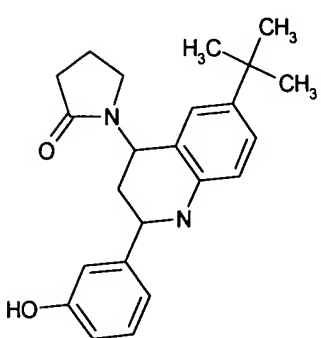
(486)



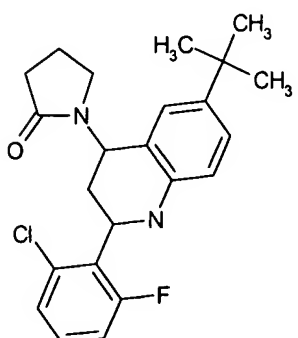
(487)



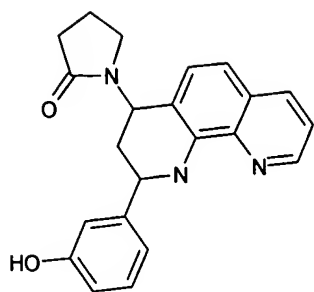
(488)



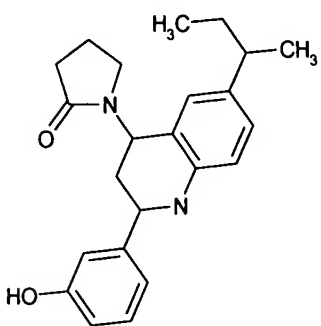
(489)



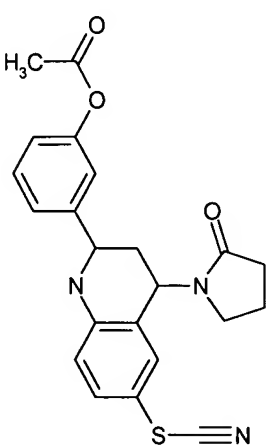
(490)



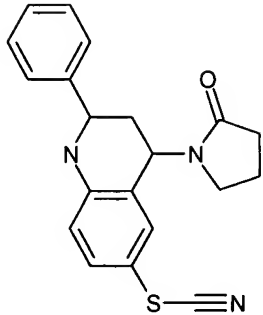
(491)



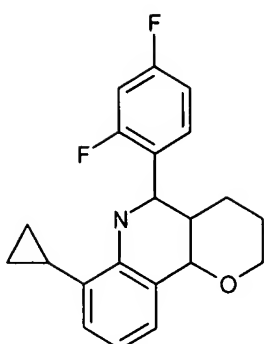
(492)



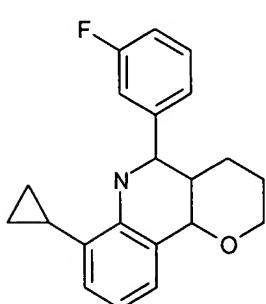
(493)



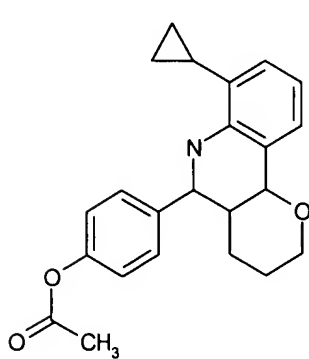
(494)



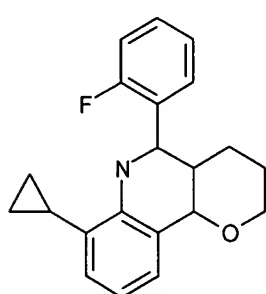
(495)



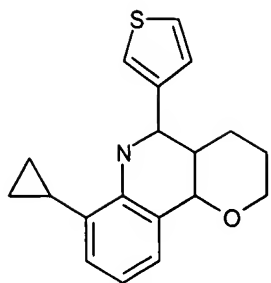
(496)



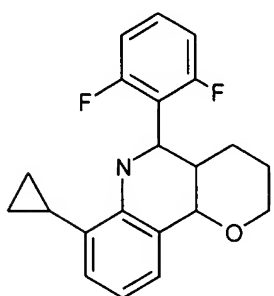
(497)



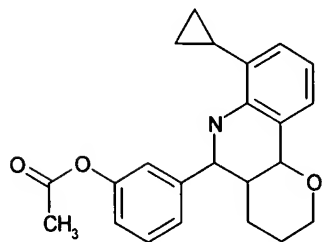
(498)



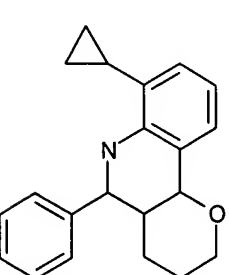
(499)



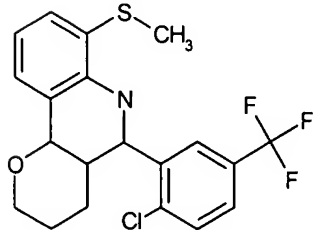
(500)



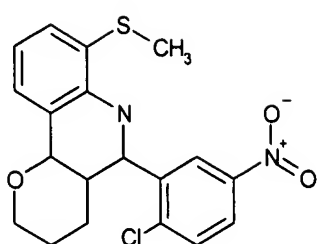
(501)



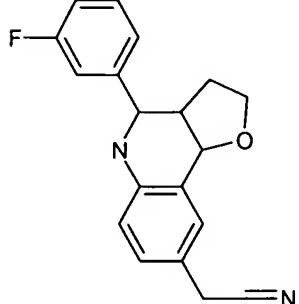
(502)



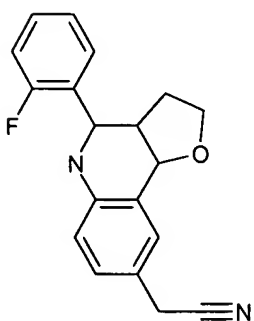
(503)



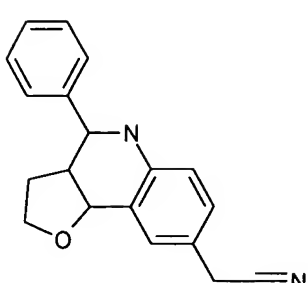
(504)



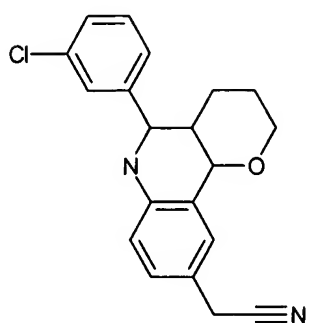
(505)



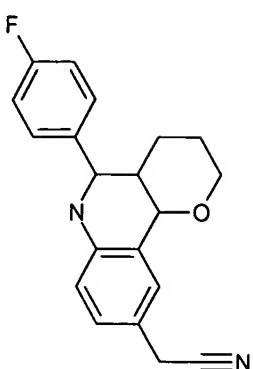
(506)



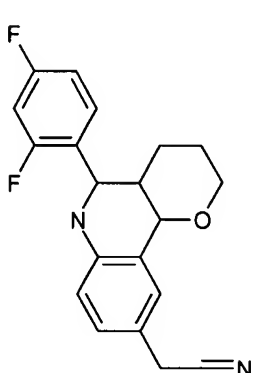
(507)



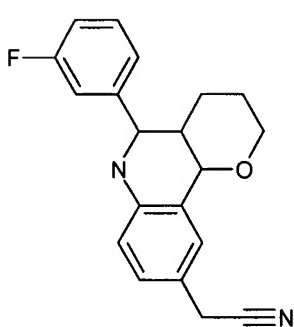
(508)



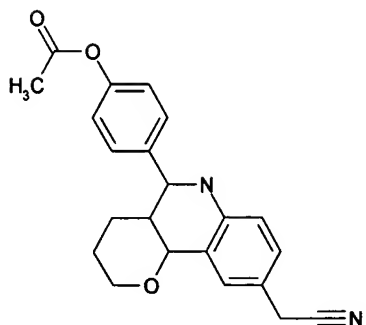
(509)



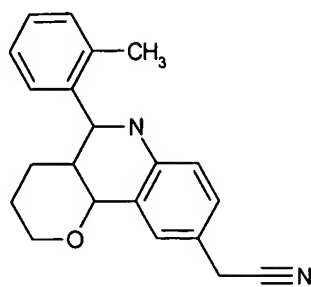
(510)



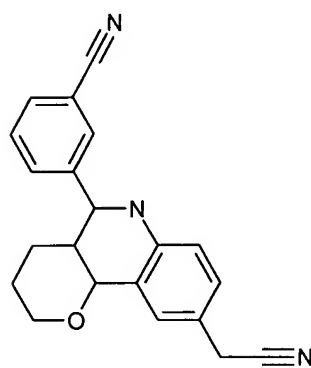
(511)



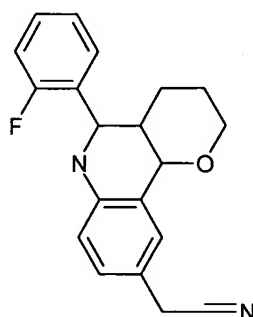
(512)



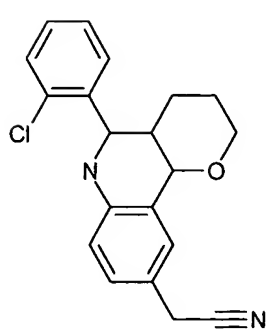
(513)



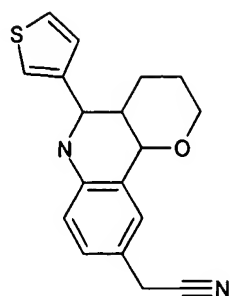
(514)



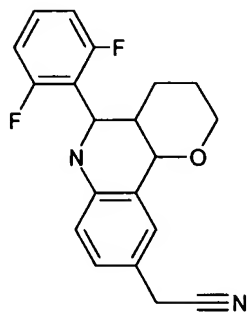
(515)



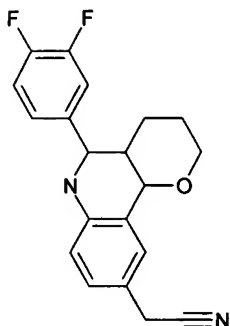
(516)



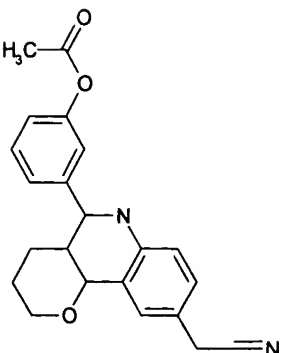
(517)



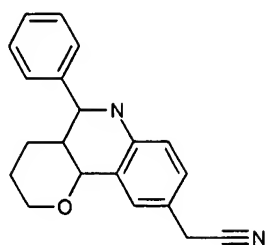
(518)



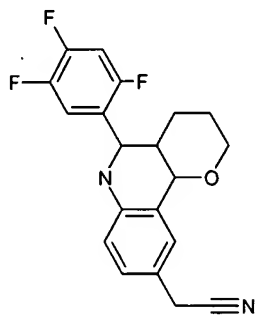
(519)



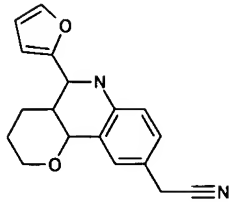
(520)



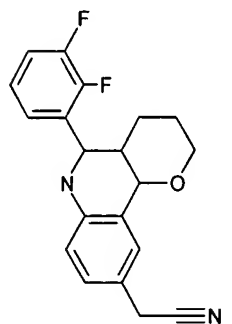
(521)



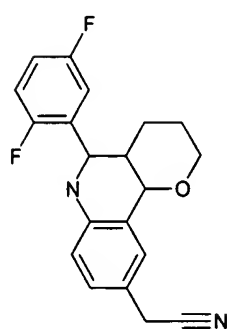
(522)



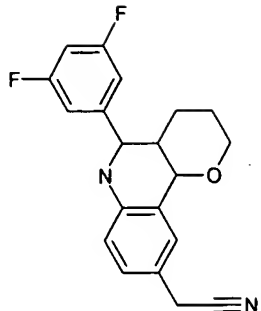
(523)



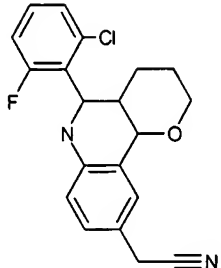
(524)



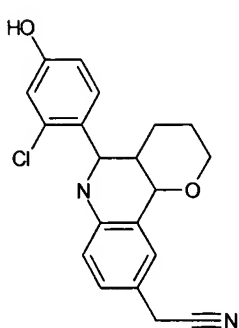
(525)



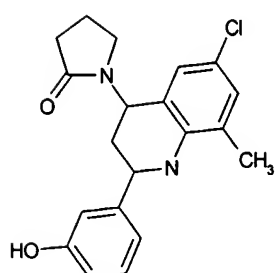
(526)



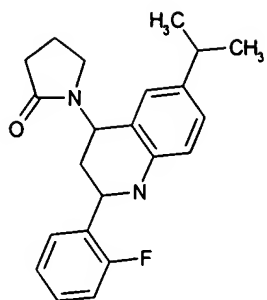
(527)



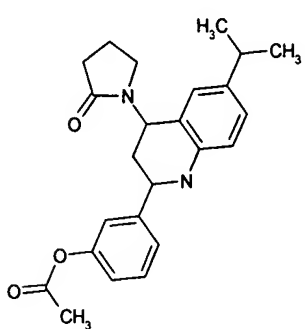
(528)



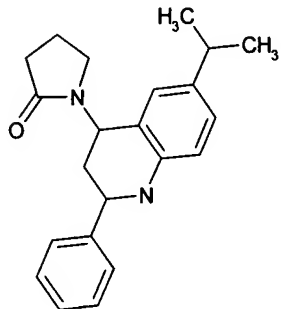
(529)



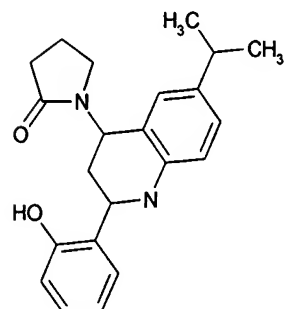
(530)



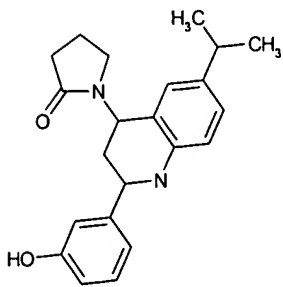
(531)



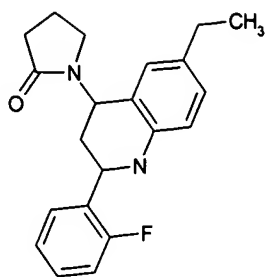
(532)



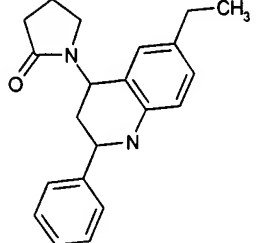
(533)



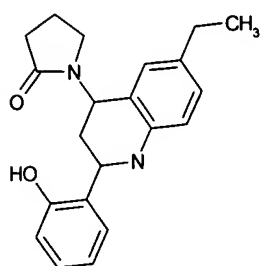
(534)



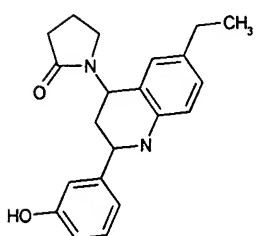
(535)



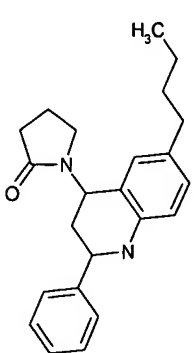
(536)



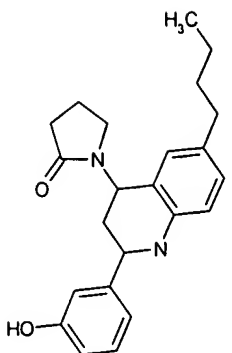
(537)



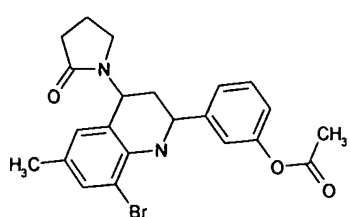
(538)



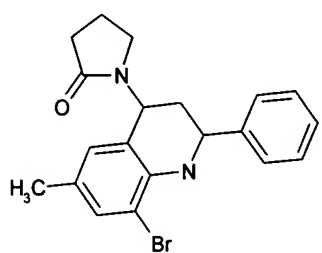
(539)



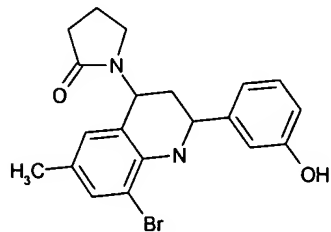
(540)



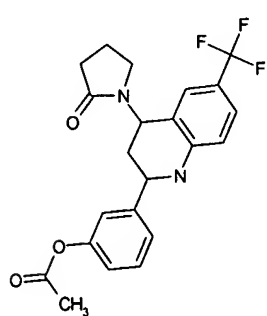
(541)



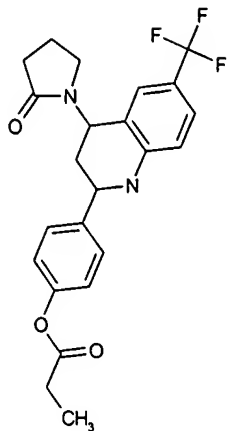
(542)



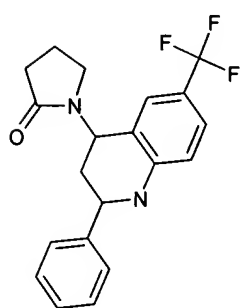
(543)



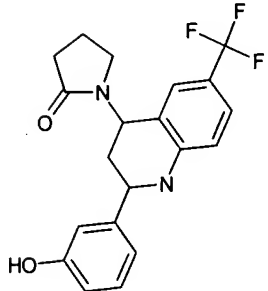
(544)



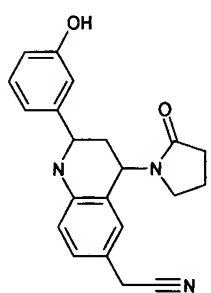
(545)



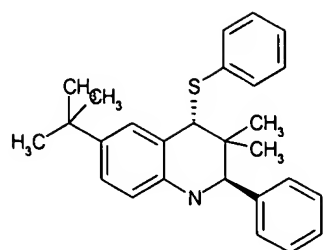
(546)



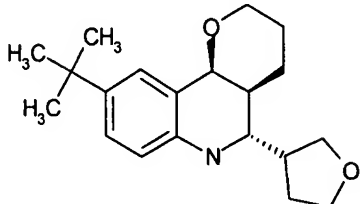
(547)



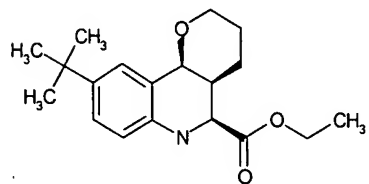
(548)



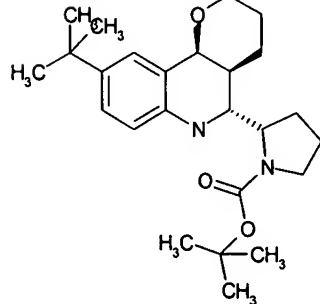
(549)



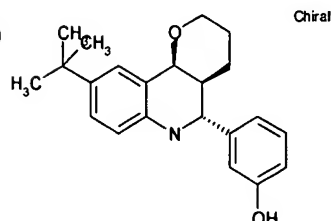
(550)



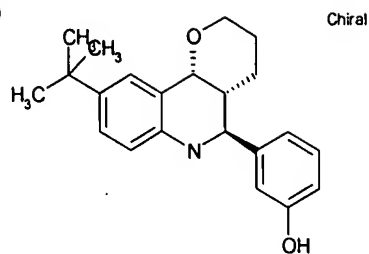
(551)



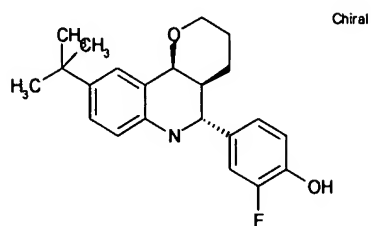
(552)



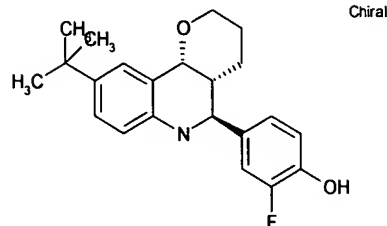
(553)



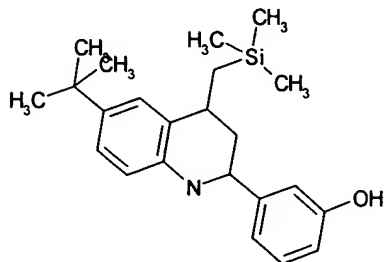
(554)



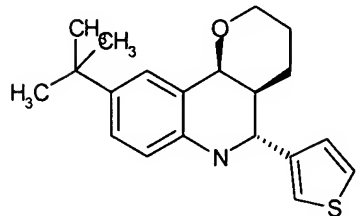
(555)



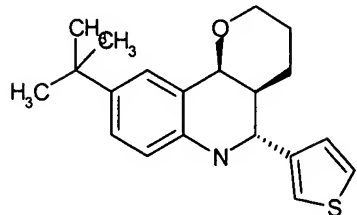
(556)



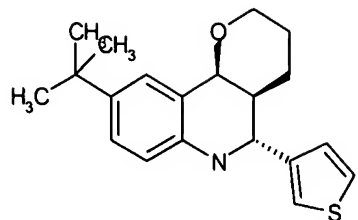
(557)



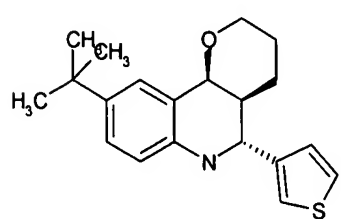
(558)



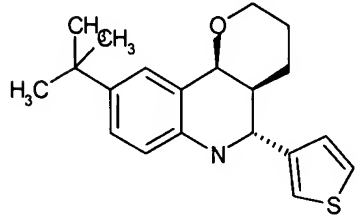
(559)



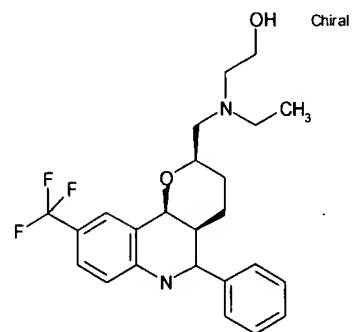
(560)



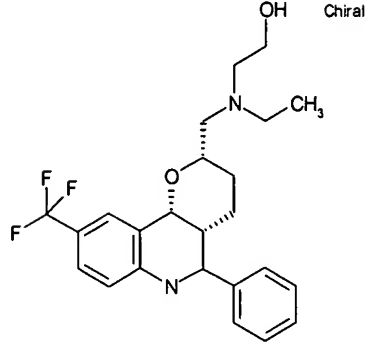
(561)



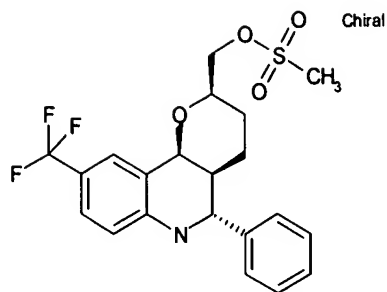
(562)



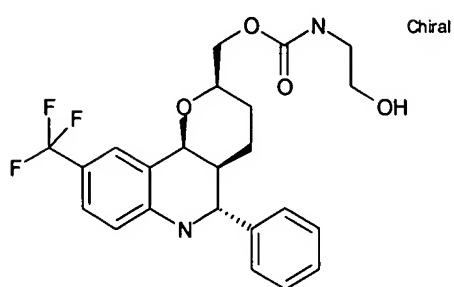
(563)



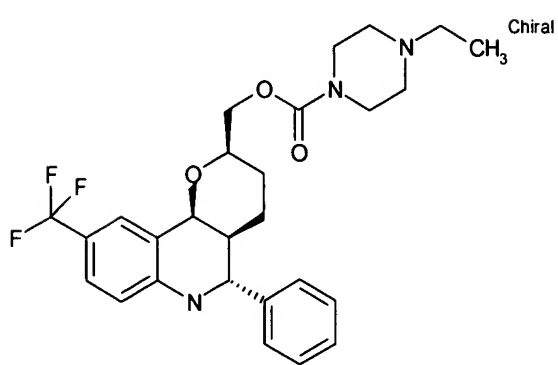
(564)



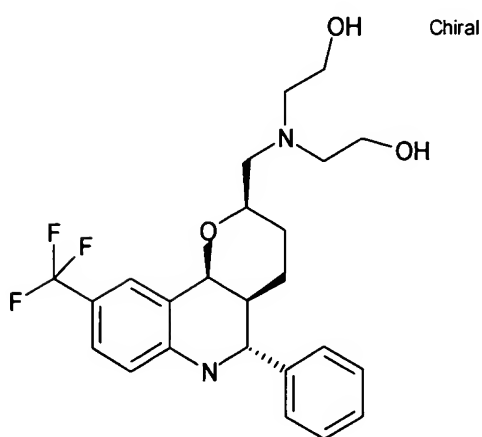
(565)



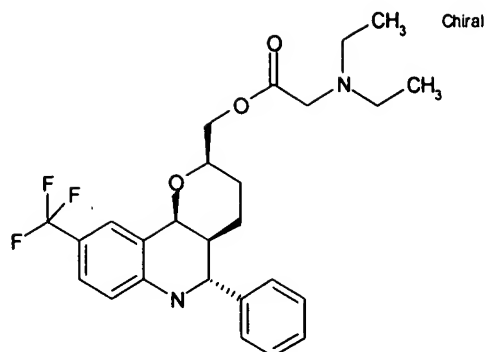
(566)



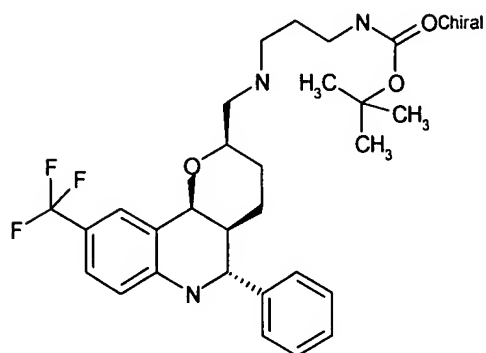
(567)



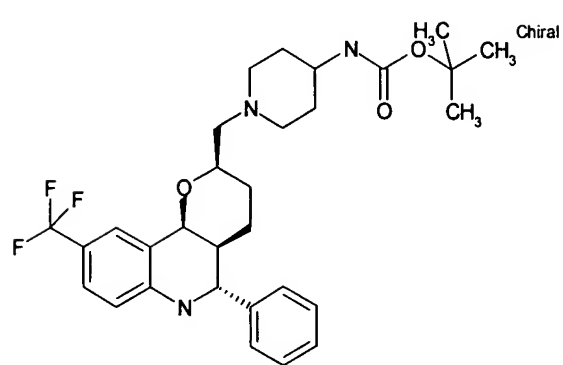
(568)



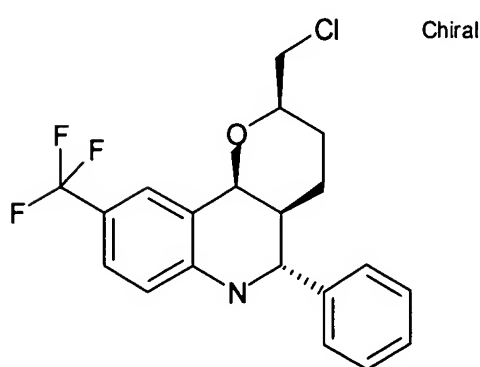
(569)



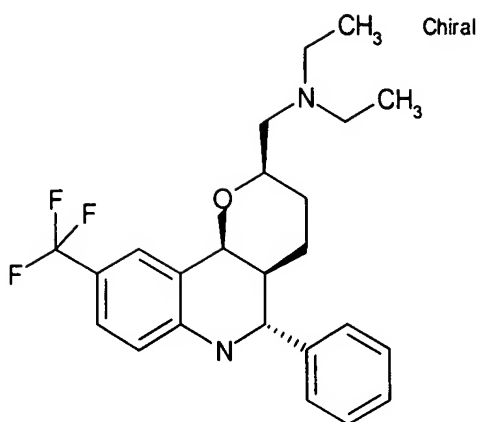
(570)



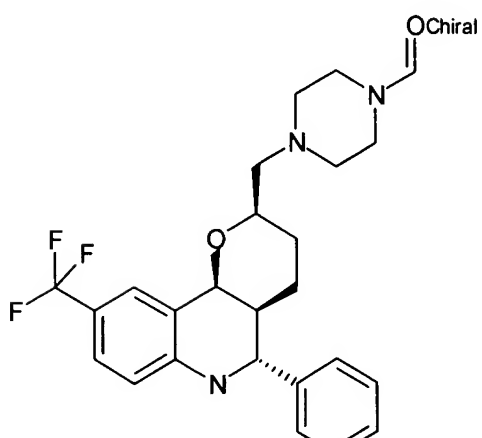
(571)



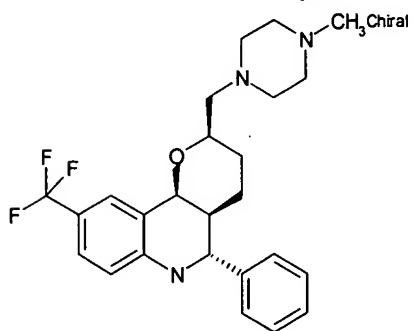
(572)



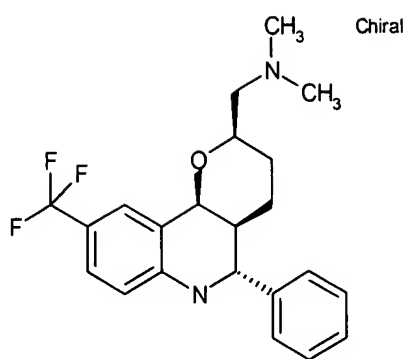
(573)



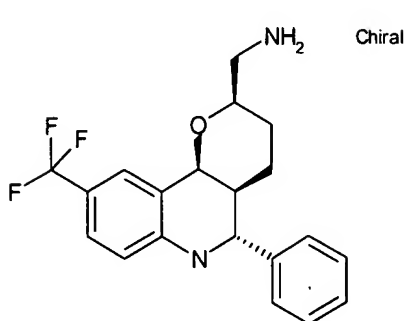
(574)



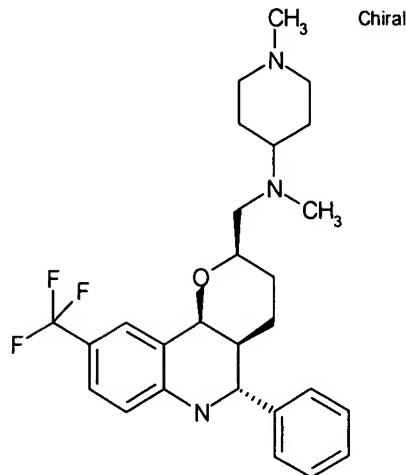
(575)



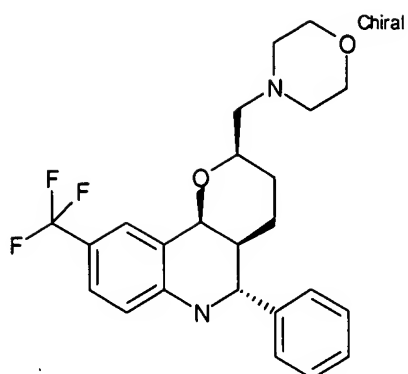
(576)



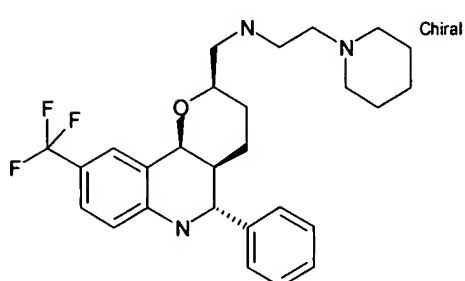
(577)



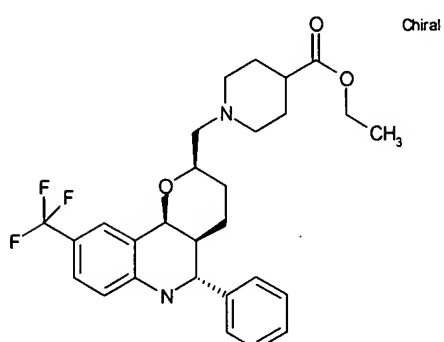
(578)



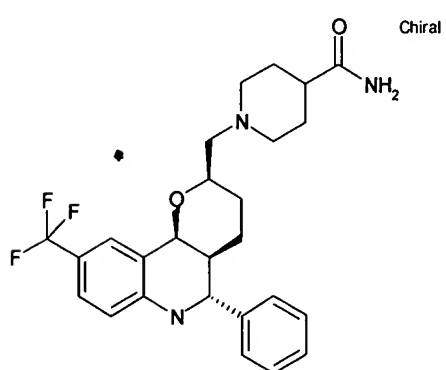
(579)



(580)

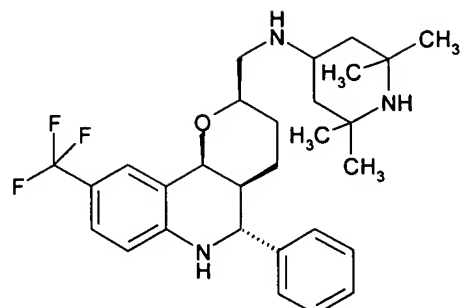


(581)



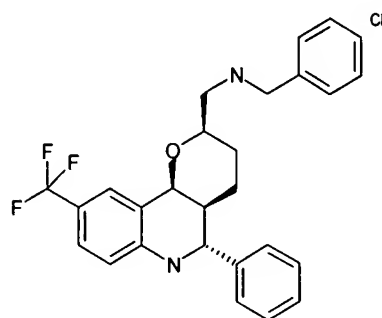
(582)

Chiral



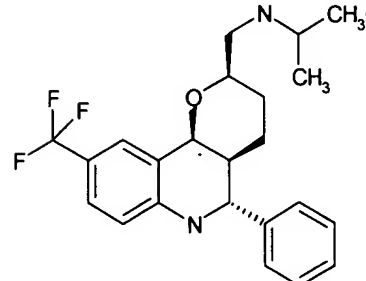
(583)

Chiral



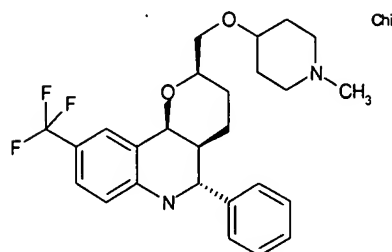
(584)

Chiral

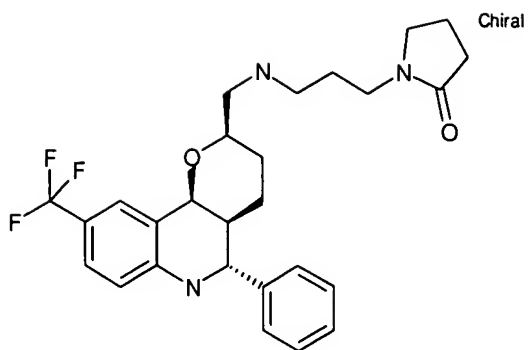


(585)

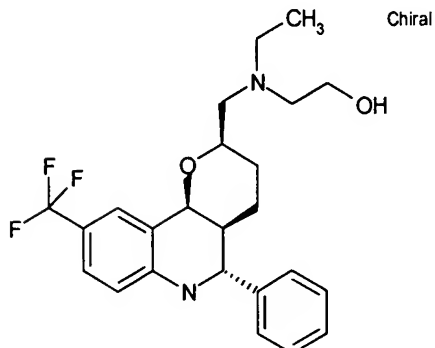
Chiral



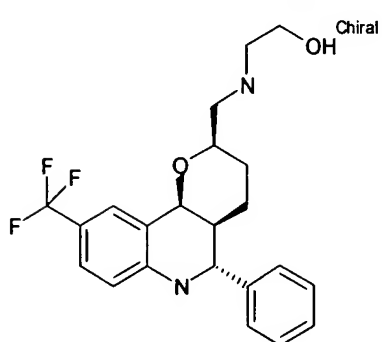
(586)



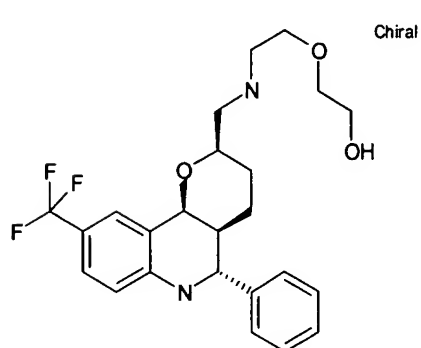
(587)



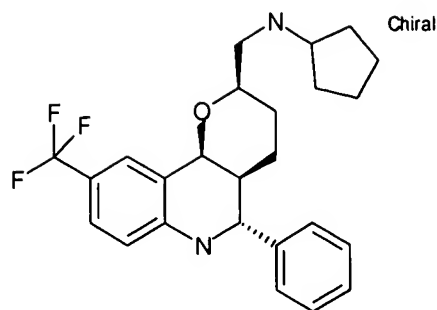
(588)



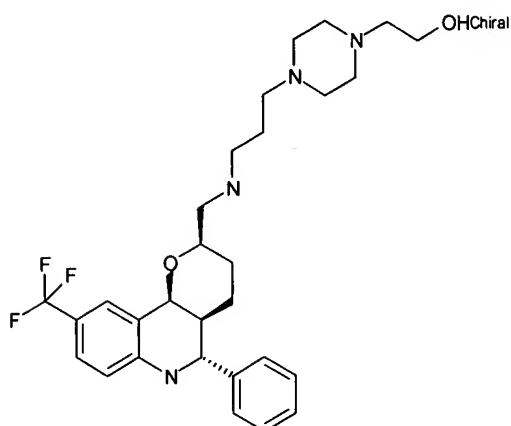
(589)



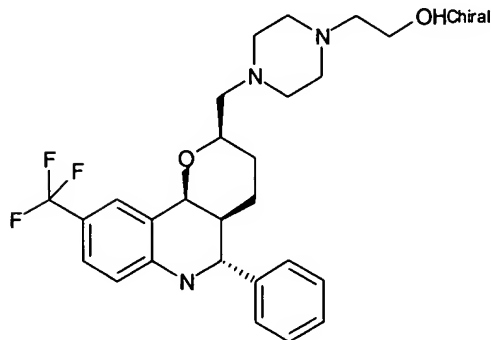
(590)



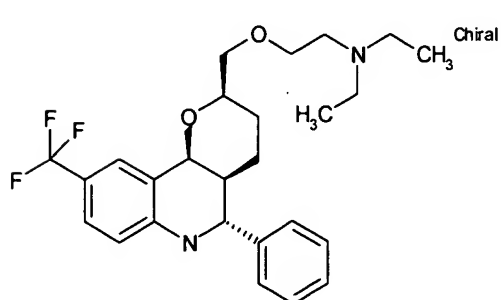
(591)



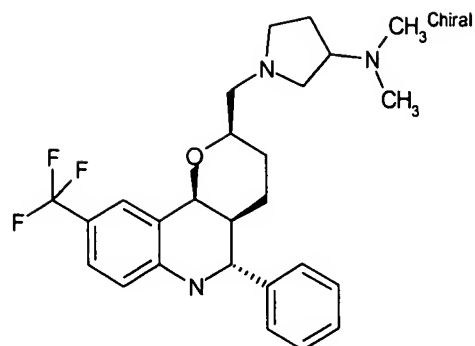
(592)



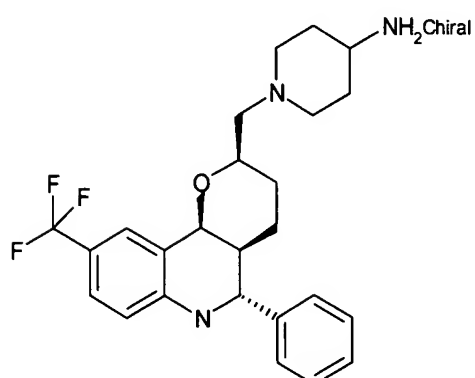
(593)



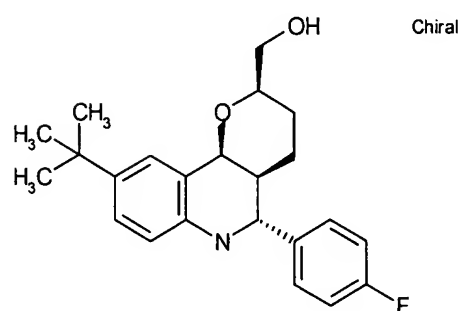
(594)



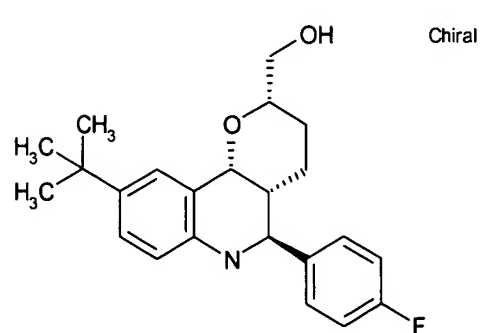
(595)



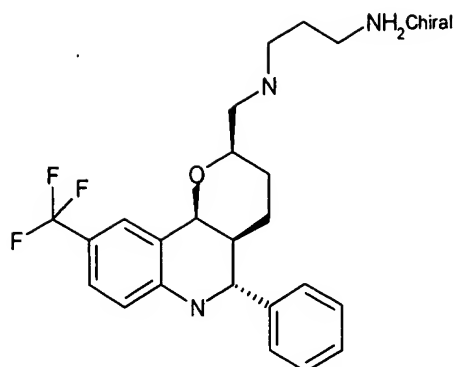
(596)



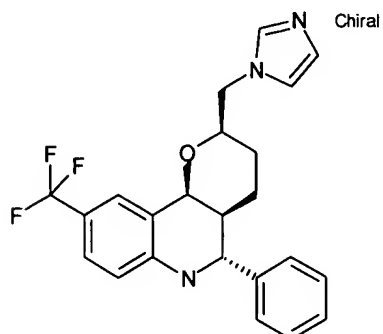
(597)



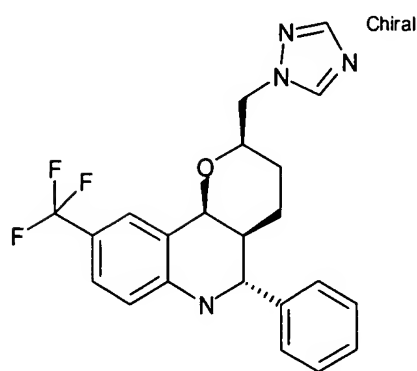
(598)



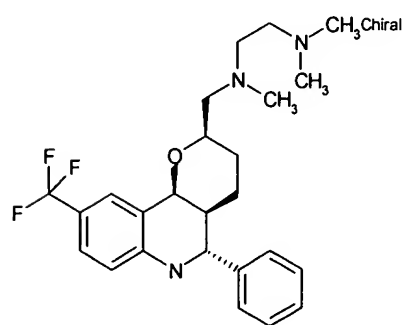
(599)



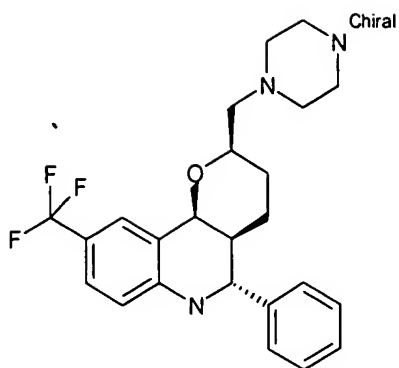
(600)



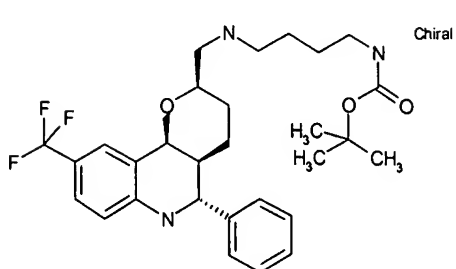
(601)



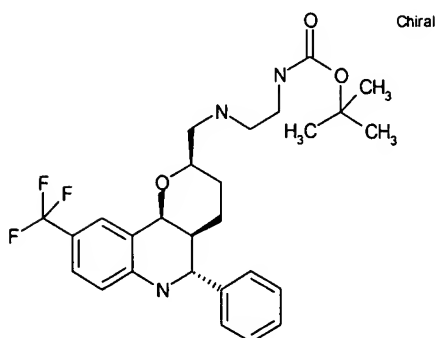
(602)



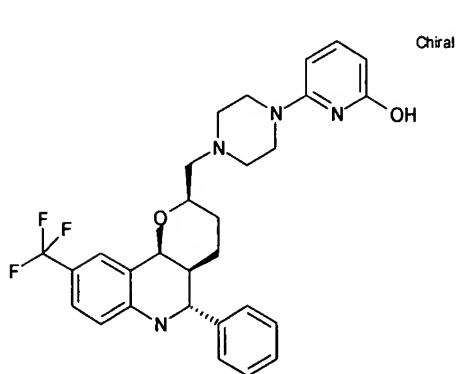
(603)



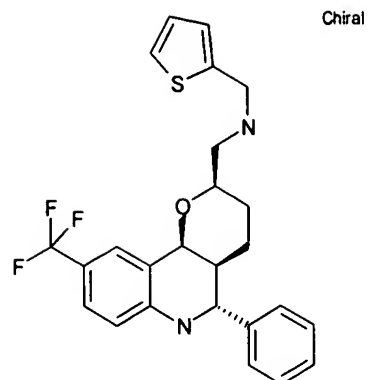
(604)



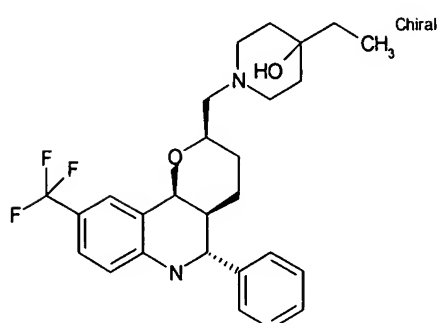
(605)



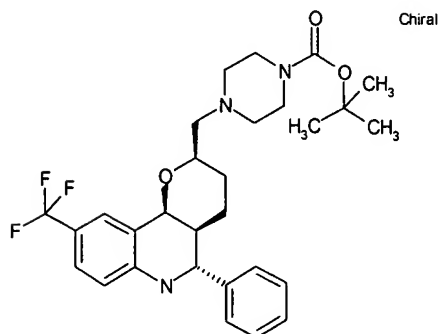
(606)



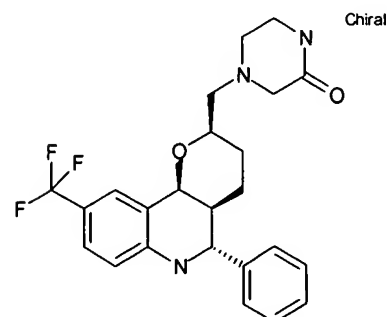
(607)



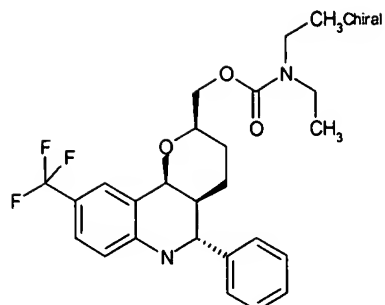
(608)



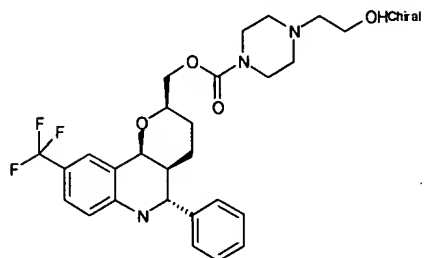
(609)



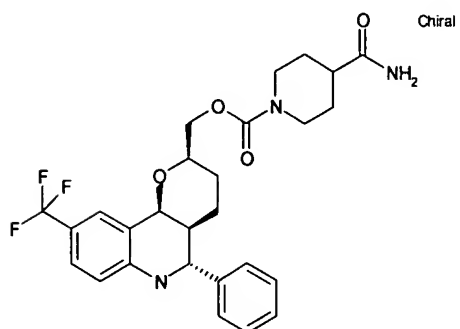
(610)



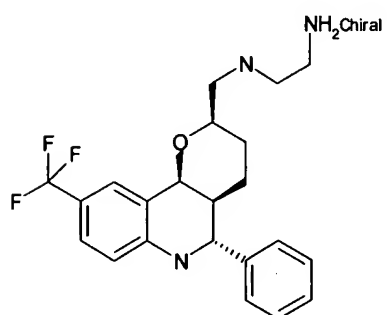
(611)



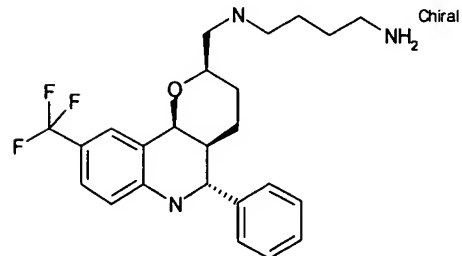
(612)

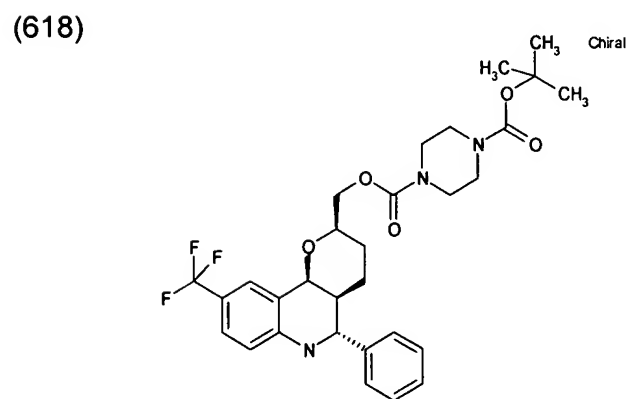
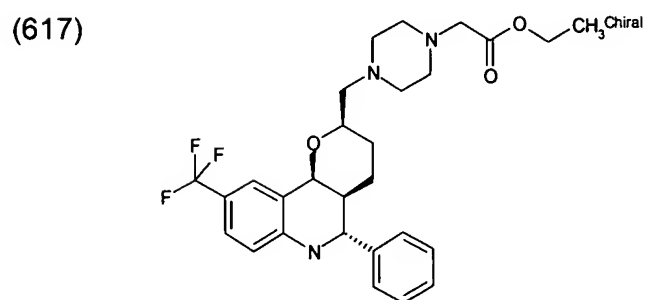
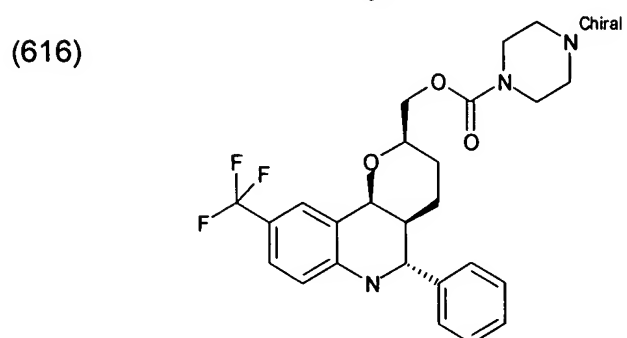
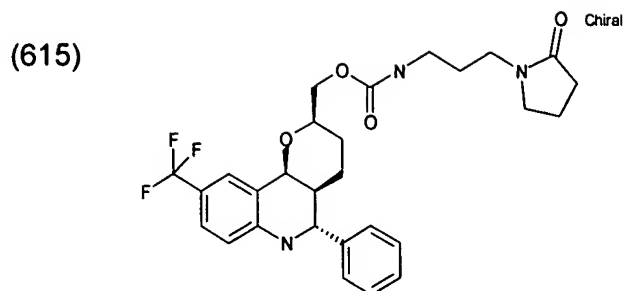


(613)

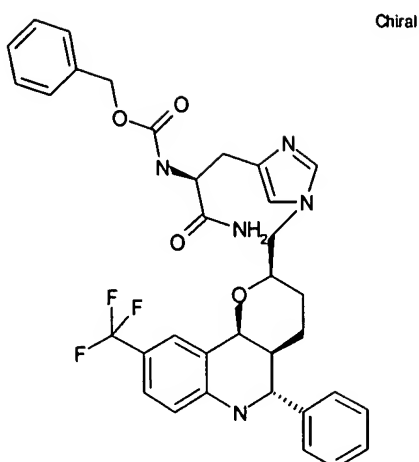


(614)



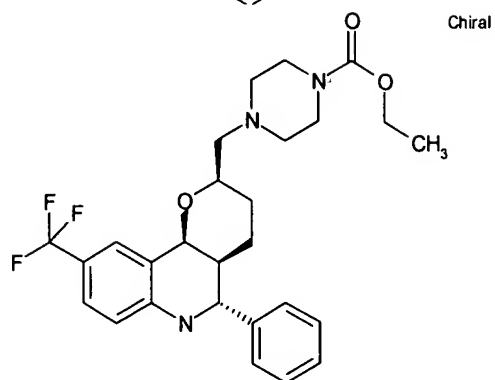


(619)



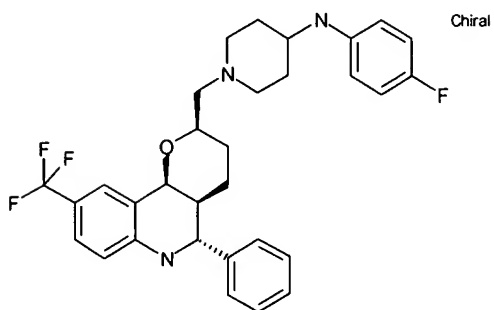
Chiral

(620)



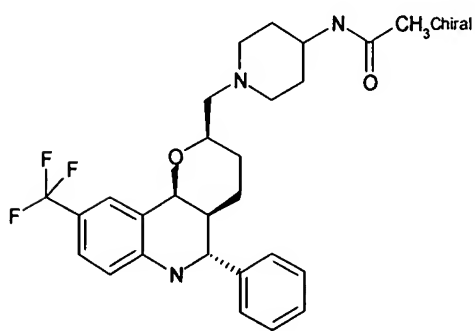
Chiral

(621)



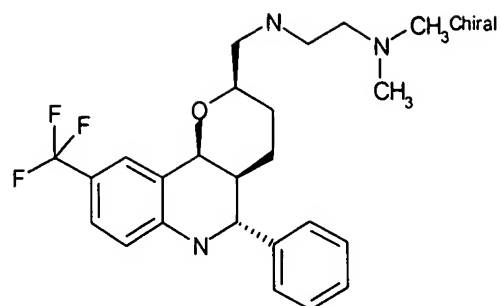
Chiral

(622)

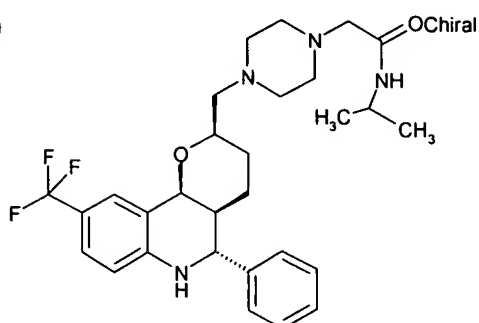


Chiral

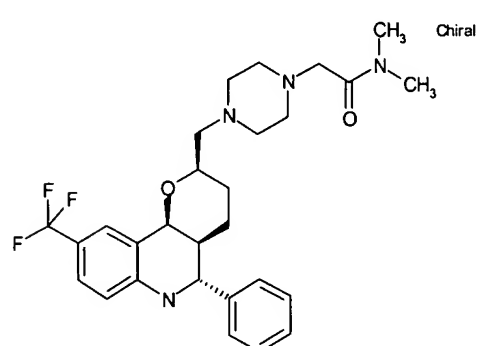
(623)



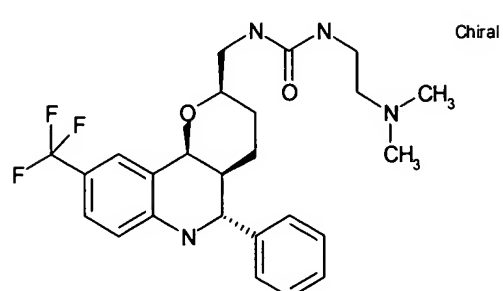
(624)

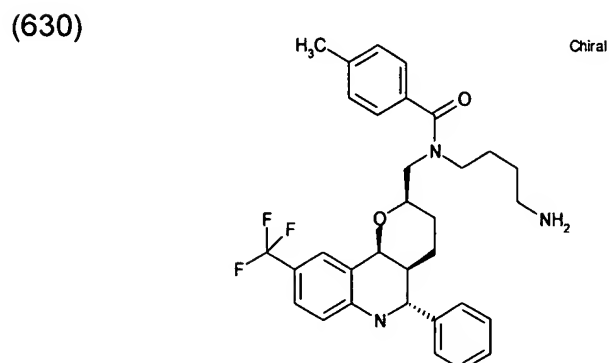
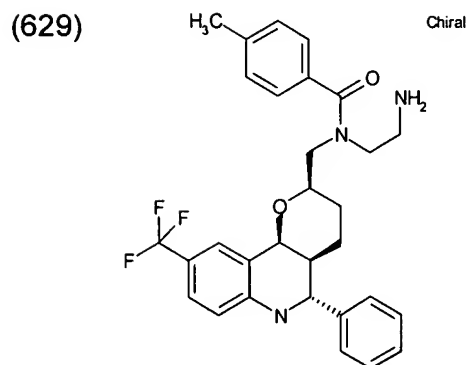
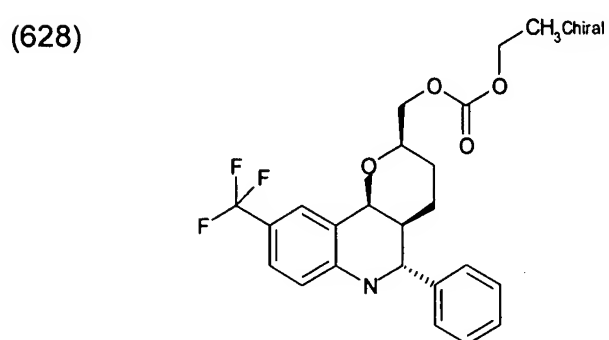
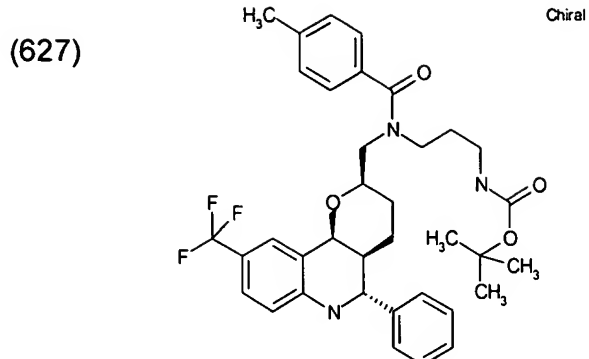


(625)

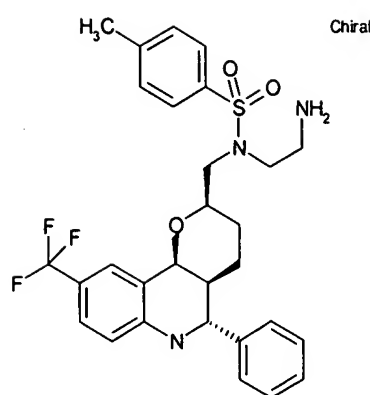


(626)

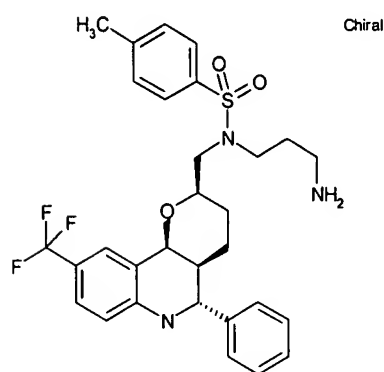




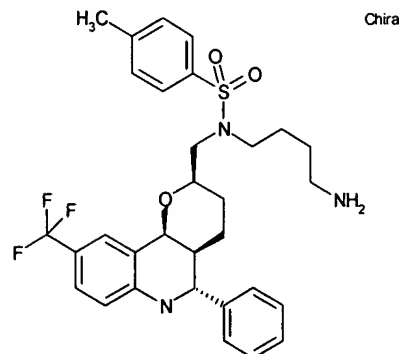
(631)



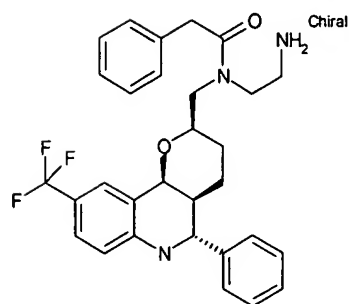
(632)



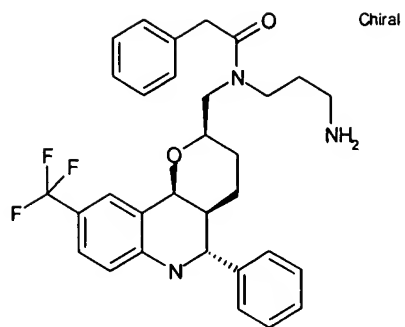
(633)



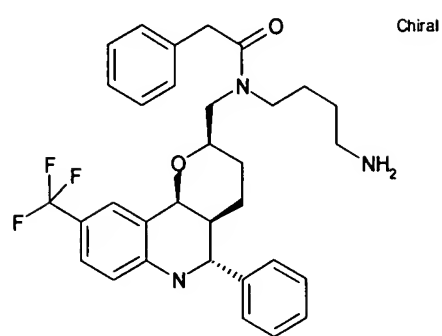
(634)



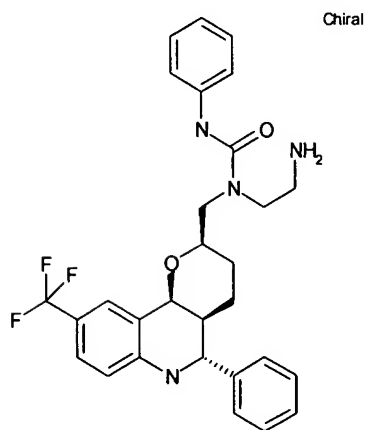
(635)



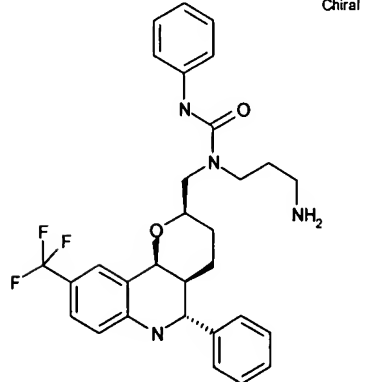
(636)



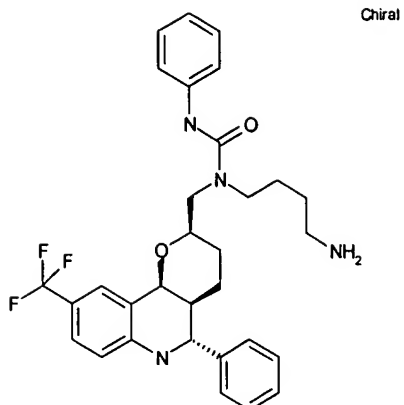
(637)



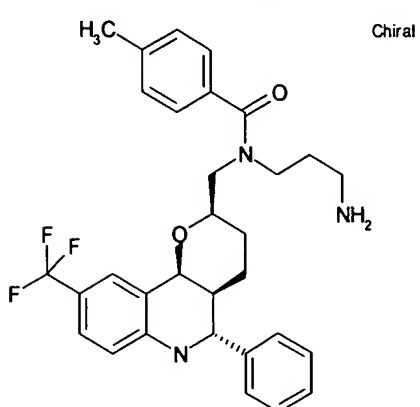
(638)



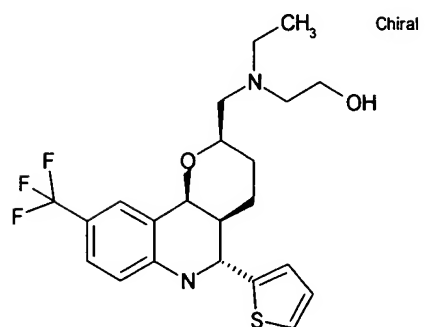
(639)



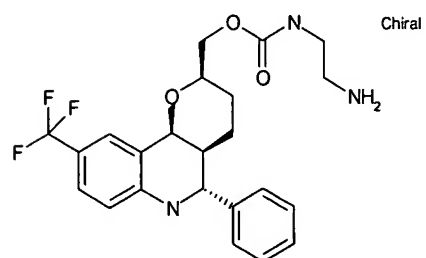
(640)



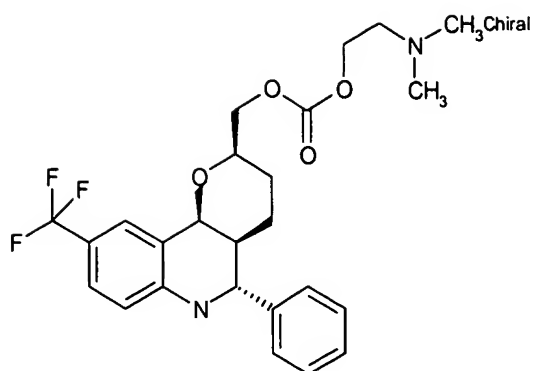
(641)



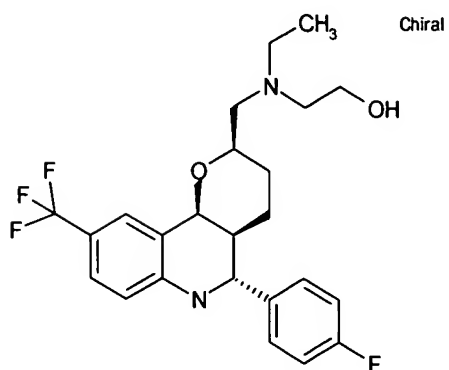
(642)



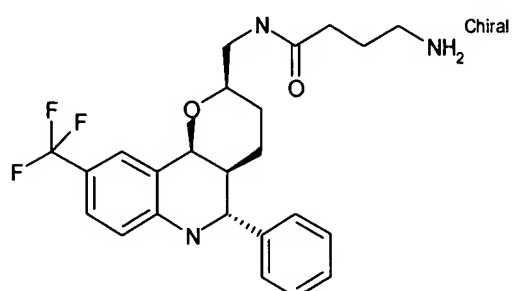
(643)



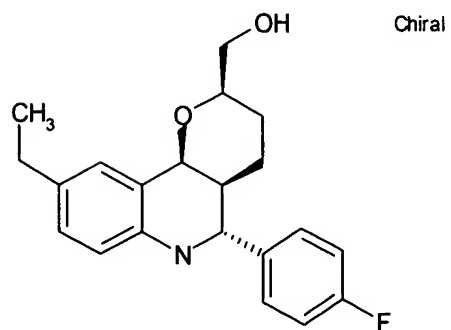
(644)



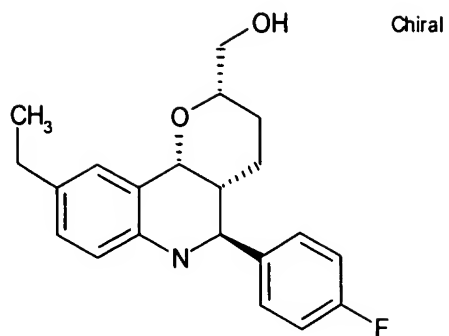
(645)



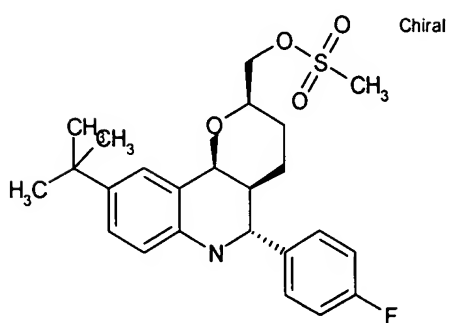
(646)



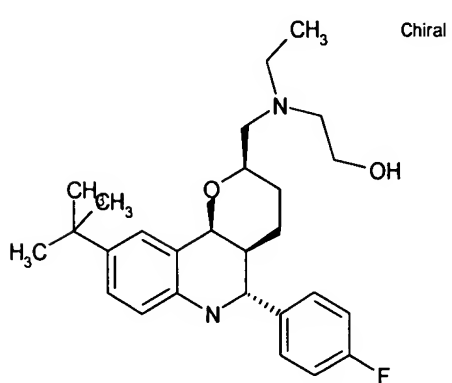
(647)



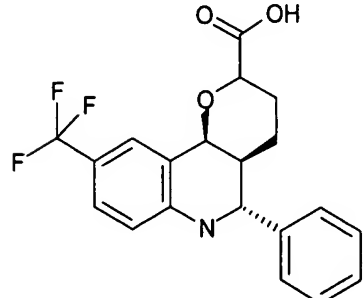
(648)



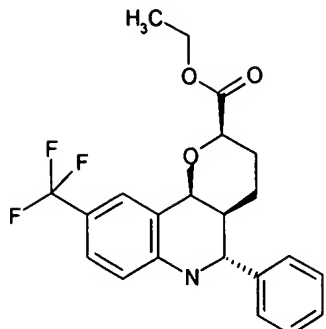
(649)



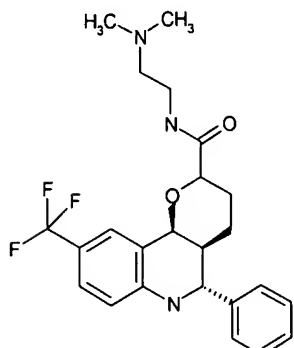
(650)



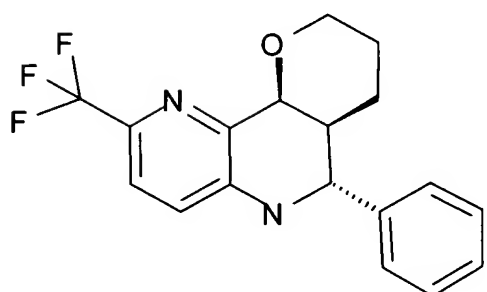
(651)



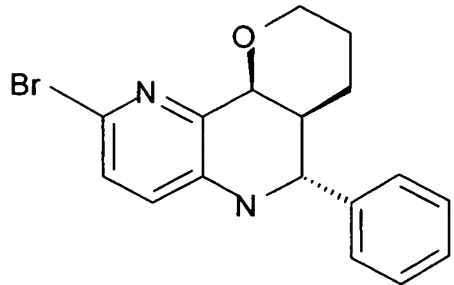
(652)



(653)



(654)



The compounds of the formula I from Examples 562 to 654 are particularly preferred here since they exhibit an increased action according to the invention.

Example A: Assay I

The efficacy of the compounds according to the invention can be determined, for example, via the Eg5 ATPase activity, which is measured via an enzymatic regeneration of the product ADP to ATP by means of pyruvate kinase (PK) and subsequent coupling to an NADH-dependent lactate dehydrogenase (LDH) reaction. The reaction can be monitored via the change in absorbance at 340 nm by coupling to the NADH-dependent LDH. The regeneration of the ATP simultaneously ensures that the substrate concentration remains constant. The change in absorbance per time unit are analysed graphically and a linear regression carried out in the visually linear region of the reaction.

Example B: Assay II

The combination of the antiprotozoic pentamidine and the inhibitors of kinesin ATPase Eg5/KSP results in increased inhibitory effects in cell proliferation tests with the colon carcinoma cell line HCT116.

Eg5 inhibitors adversely affect the ATPase activity and inhibit the course of the cell cycle owing to an error in the separation of the spindle poles.

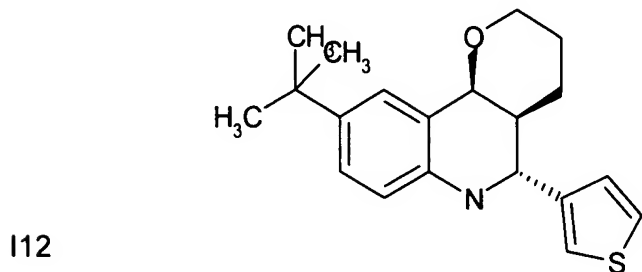
The determination of the efficacy of the compounds of the formula I according to the invention in combination with compounds of the formula V and/or medicaments from Table I can be demonstrated as follows in combination assays:

10^3 to 10^4 cells of a defined cell line (HCT116, Colo 205, MDA-MB 231, etc.) are sown into each well of a 96-well microtitre plate and cultivated overnight under standard conditions. For the substances of the combination

to be tested, 10-50 mM stock solutions in DMSO were prepared. Dilution series (generally 3-fold dilution steps) of the individual substances were combined with one another in the form of a pipetting scheme (see scheme below), while maintaining a DMSO final concentration of 0.5% (v/v). Next morning, the substance mixtures were added to the cells, which were incubated under culture conditions for a further 48 hours. At the end of the cultivation, Crystal Violet staining of the cells was carried out. After extraction of the Crystal Violet from the fixed cells, the absorption at 550 nm was measured spectrophotometrically. It can be used as a quantitative measure of the adherent cells present.

Scheme

The following results give by way of example indications of the inhibition of the cell cycle by compound I12 according to the invention



Monasterol and pentamidine in individual experiments:

- HCT116 cells were treated with the given concentrations of the compounds for 24 hrs, processed for the propidium iodide analysis with a BD-FACS scan in order to determine the DNA content of the treated cell populations. The percentage of cells in the G0/G1 phase (unreplicated DNA), S phase (DNA replication phase) and G2/M phase (duplicated DNA before mitosis) is shown. 10,000 events were counted for each concentration. Striking changes in the G2/M enrichment have been highlighted.

I12 (Eg5 inhibitor according to the invention):

μM	G1	S	G2/M
0	36	43	21
0.0032	36	44	21
0.016	36	43	21
0.08	35	42	23
0.4	7	7	86
2	0	15	85

Monastrol (Eg5 inhibitor from the prior art)

μM	G1	S	G2/M
0	41	38	21
0.08	35	39	25
0.4	34	38	28
2	33	42	25
10	34	39	27
50	14	33	52

Pentamidine isothionate

μM	G1	S	G2/M
0	36	42	22
0.016	36	42	22
0.08	35	44	21
0.4	33	45	22
2	32	44	24
10	32	32	32

The following results give by way of example indications of the inhibition of the cell cycle by compound I12 according to the invention or monasterol and pentamidine in combination experiments:

Combination therapy: I12 plus pentamidine

HCT116 cells were incubated overnight at 37°C in 96-well plates at a density of 2500 or 5000 cells/well. On the following day, a concentration matrix of pentamidine and the compounds indicated was added, and the cells were incubated for a further 48 hrs. The cells were then stained with Crystal Violet, and the absorption was measured after extraction of the dye. The absorption value reflects the number of cells in the well.

Pipetting scheme:

<--Pentamidine--						
[μM]	32	16	8	4	2	0
[M2] [μM]	0.4					
	0.2					
	0.1					
	0.05					
	0.02					
	5					
	0					

Results:

Upper matrix: OD550 nm background corrected

Central matrix: % of control (the 100% value stands for vehicle-treated cells minus background signal)

Lower field: -fold inhibition values (i.e. the 100%/x% value stands for a -fold inhibition of proliferation).

Table 2

Pentamidine [μ M]	32	16	8	4	2	0
0.4	0.1	0.2	0.3	0.3	0.4	0.4
0.2	0.2	0.4	0.7	1.1	1.0	1.1
0.1	0.2	0.4	0.7	1.0	1.1	1.1
0.05	0.2	0.4	0.7	1.1	1.1	1.2
0.025	0.2	0.4	0.7	1.1	1.1	1.1
0	0.2	0.4	0.6	1.0	1.0	1.1
[I2 [μ M]]						
Pentamidine [μ M]	32	16	8	4	2	0
0.4	9	14	21	25	32	31
0.2	20	33	55	88	82	90
0.1	17	34	61	86	93	90
0.05	16	31	60	90	93	98
0.025	17	31	58	92	91	91
0	20	30	50	85	86	90
[I2 [μ M]]						
Pentamidine [μ M]	32	16	8	4	2	0
0.4	11	7	5	4	3	3
0.2	5	3	2	1	1	1
0.1	6	3	2	1	1	1
0.05	6	3	2	1	1	1
0.025	6	3	2	1	1	1
0	5	3	2	1	1	1
[I2 [μ M]]						

Graphical representation of the data:

The data have been represented in FIG 1 as concentration curves (each line illustrates one curve at a given pentamidine concentration)

Combination therapy: monastrol plus pentamidine**Pipetting scheme:**

		Pentamidine					
		32	16	8	4	2	0
Monastrol ↑	50						
	25						
	12.5						
	6.25						
	0						

Results:

Upper matrix: OD550 nm background corrected

Central matrix: % of control (the 100% value stands for vehicle-treated cells minus background signal)

Lower field: -fold inhibition values (i.e. the 100%/x% value stands for a -fold inhibition of proliferation).

Table 3

Pentamidine [µM]	16	8	4	2	0
50	0.02	0.12	0.19	0.23	0.14
25	0.07	0.15	0.24	0.34	0.43
12.5	0.07	0.18	0.57	0.73	0.51
6.125	0.11	0.34	0.57	1.08	0.68
0	0.15	0.45	0.82	1.01	0.54
Monastrol					
Pentamidine [µM]	16	8	4	2	0
50	3	15	24	29	17
25	9	19	29	42	53
12.5	8	22	71	89	62
6.125	13	42	70	132	84
0	18	55	101	124	67
Monastrol					
Pentamidine [µM]	16	8	4	2	0
50	34	7	4	3	6
25	12	5	3	2	2
12.5	12	5	1	1	2
6.125	7	2	1	1	1
0	6	2	1	1	1
Monastrol					

Graphical representation of the data:

The data have been represented in FIG 2 as concentration curves (each line illustrates one curve at a given pentamidine concentration)

The following examples relate to medicaments:

Example C: Injection vials

A solution of 100 g of an active ingredient of the formula I and 5 g of disodium hydrogenphosphate in 3 l of bidistilled water is adjusted to pH 6.5

using 2 N hydrochloric acid, sterile filtered, transferred into injection vials, lyophilised under sterile conditions and sealed under sterile conditions. Each injection vial contains 5 mg of active ingredient.

Example D: Suppositories

A mixture of 20 g of an active ingredient of the formula I with 100 g of soya lecithin and 1400 g of cocoa butter is melted, poured into moulds and allowed to cool. Each suppository contains 20 mg of active ingredient.

Example E: Solution

A solution is prepared from 1 g of an active ingredient of the formula I, 9.38 g of $\text{NaH}_2\text{PO}_4 \cdot 2 \text{ H}_2\text{O}$, 28.48 g of $\text{Na}_2\text{HPO}_4 \cdot 12 \text{ H}_2\text{O}$ and 0.1 g of benzalkonium chloride in 940 ml of bidistilled water. The pH is adjusted to 6.8, and the solution is made up to 1 l and sterilised by irradiation. This solution can be used in the form of eye drops.

Example F: Ointment

500 mg of an active ingredient of the formula I are mixed with 99.5 g of Vaseline under aseptic conditions.

Example G: Tablets

A mixture of 1 kg of active ingredient of the formula I, 4 kg of lactose, 1.2 kg of potato starch, 0.2 kg of talc and 0.1 kg of magnesium stearate is pressed in a conventional manner to give tablets in such a way that each tablet contains 10 mg of active ingredient.

Example H: Dragees

Tablets are pressed analogously to Example E and subsequently coated in a conventional manner with a coating of sucrose, potato starch, talc, tragacanth and dye.

Example I: Capsules

2 kg of active ingredient of the formula I are introduced into hard gelatine capsules in a conventional manner in such a way that each capsule contains 20 mg of the active ingredient.

Example J: Ampoules

A solution of 1 kg of active ingredient of the formula I in 60 l of bidistilled water is sterile filtered, transferred into ampoules, lyophilised under sterile conditions and sealed under sterile conditions. Each ampoule contains 10 mg of active ingredient.

Description of the figures**FIG 1**

Graphical representation of the data from Table 2

The data have been represented as concentration curves of the substance of the formula I12. Each line illustrates one curve at a given pentamidine concentration.

FIG 2

Graphical representation of the data from Table 3

The data have been represented as concentration curves of the substance monastrol. Each line illustrates one curve at a given pentamidine concentration.

FIG 3 to FIG 25

NMR spectra of representative compounds of the formula I